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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREAT (1917)					
(51) International Patent Classification 6:		(11) International Publication Number: WO 95/19369			
C07H 21/02, 21/04, C12Q 1/68, G01N 33/53, C12P 21/00, C12N 15/63, 15/85, A61K 48/00	A1	(43) International Publication Date: 20 July 1995 (20.07.95)			
(21) International Application Number: PCT/US95/00608 (22) International Filing Date: 17 January 1995 (17.01.95)		(81) Designated States: AU, CA, JP, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).			
(30) Priority Data: 08/182,961 08/373,799 14 January 1994 (14.01.94) 17 January 1995 (17.01.95)		Published With international search report. With amended claims and statement.			
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(54) Title: METHOD FOR DETECTION AND TREATMENT OF BREAST CANCER

(57) Abstract

The present invention provides a method of detecting and diagnosing pre-invasive breast cancer by identifying differentially expressed genes in early, pre-invasive breast cancer tissue. Differentially expressed genes can be used as genetic markers to indicate the presence of pre-invasive cancerous tissues. Microscopically directed tissue sampling techniques combined with differential display or differential screening of cDNA libraries are used to determine differential expression of genes in the early stages of breast cancer. Differential expression of genes in pre-invasive breast cancer tissue is confirmed by RT-PCR, nuclease protection assays and in-situ hybridization of ductal carcinoma in situ tissue RNA and control tissue RNA. The present invention also provides a method of screening for compounds that induce expression of the BRCA1 gene, whose product negatively regulates cell growth in both normal and malignant mammary epithlial cells. The present invention also relates to gene therapy method using this gene.

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DESCRIPTION

"METHOD FOR DETECTION AND TREATMENT OF BREAST CANCER" TECHNICAL FIELD

The present invention relates generally to methods of detection and diagnosis of breast cancer and more particularly to a diagnostic method which relies on the identification of marker genes expressed in pre-invasive cancers by microscopically-directed cloning. Furthermore, this invention concerns the prevention, detection, and diagnosis of breast cancer by addressing the molecular events which occur during the earliest alterations in breast tissue.

The present invention also relates generally to methods of treatment of breast cancer, and more particularly to gene therapy methods and methods for screening compounds that induce expression of the BRCA1 gene product.

BACKGROUND ART

It will be appreciated by those skilled in the art that there exists a need for a more sensitive and less invasive method of early detection and diagnosis of breast cancer than those methods currently in use. Breast cancer presents inherent difficulties in regard to the ease with which it is detected and diagnosed. This is in contrast to detection of some other common cancers, including skin and cervical cancers, the latter of which is based on cytomorphologic screening techniques.

There have been several attempts to develop improved methods of breast cancer detection and diagnosis. In the attempts to improve methods of detection and diagnosis of breast cancer, numerous studies have scarched for oncogene mutations, gene amplification, and loss of heterozygosity in invasive breast cancer (Callahan, et al., 1992; Cheickh, et al., 1992; Chen, et al., 1992; and, Lippman, et al, 1990). However, few studies of breast cancer have analyzed gene mutations and/or altered gene expression in ductal carcinoma in situ (DCIS). Investigators have demonstrated high levels of p53 protein in 13-40% of DCIS lesions employing a monoclonal antibody to p53, and subsequent sequencing demonstrated mutations in several cases (Poller et al, 1992). The neu/erbB2 gene appears to be amplified in a subset of DCIS lesions (Allred et al, 1992; Maguire et al, 1992). Histologic analysis of DCIS cases suggests that mutations and altered gene expression events, as well as changes in chromatin and

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DNA content, occur predominantly in comedo DCIS (Böcker et al, 1992; Killeen et al, 1991; and, Komitowski et al, 1990), which has a rapid rate of local invasion and progression to metastasis. Thus, there are presently no reliable marker genes for non-comedo DCIS (NCDCIS, hereafter).

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Cancer in humans appears to be a multi-step process which involves progression from pre-malignant to malignant to metastatic disease which ultimately kills the patient. Epidemiologic studies in humans have established that certain pathologic conditions are "pre-malignant" because they are associated with increased risk of malignancy. There is precedent for detecting and eliminating pre-invasive lesions as a cancer prevention strategy: dysplasia and carcinoma in-situ of the uterine cervix are examples of pre-malignancies which have been successfully employed in the prevention of cervical cancer by cytologic screening methods. Unfortunately, because the breast cannot be sampled as readily as cervix, the development of screening methods for breast pre-malignancy involves more complex approaches than cytomorphologic screening now currently employed to detect cervical cancer.

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Pre-malignant breast disease is also characterized by an apparent morphological progression from atypical hyperplasias, to carcinoma in-situ (pre-invasive cancer) to invasive cancer which ultimately spreads and metastasizes resulting in the death of the patient. Careful histologic examination of breast biopsies has demonstrated intermediate stages which have acquired some of these characteristics but not others. Detailed epidemiological studies have established that different morphologic lesions progress at different rates, varying from atypical hyperplasia (with a low risk) to comedo ductal carcinoma-in-situ which progresses to invasive cancer in a high percentage of patients (London et al, 1991; Page et al, 1982; Page et al, 1985; Page et al, 1991; and Page et al, 1978). Family history is also an important risk factor in the development of breast cancer and increases the relative risk of these pre-malignant lesions (Dupont et al, 1985; Dupont et al, 1993; and, London et al, 1991). Of particular interest is non-comedo carcinoma-in-situ which is associated with a greater than ten-fold increased relative risk of breast cancer compared to control groups (Ottesen et al, 1992; Page et al, 1982). Two other reasons besides an increased relative risk support the concept that DCIS is pre-malignant: 1) When breast cancer occurs in

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these patients it regularly occurs in the same region of the same breast where the DCIS was found; and 2) DCIS is frequently present in tissue adjacent to invasive breast cancer (Ottesen et al, 1992; Schwartz et al, 1992). For these reasons DCIS very likely represents a rate-limiting step in the development of invasive breast cancer in women.

DCIS (sometimes called intraductal carcinoma) is a group of lesions in which the cells have grown to completely fill the duct with patterns similar to invasive cancer, but do not invade outside the duct or show metastases at presentation. DCIS occurs in two forms: comedo DCIS and non-comedo DCIS. Comedo DCIS is often a grossly palpable lesion which was probably considered "cancer" in the 19th and early 20th century and progresses to cancer (without definitive therapy) in at least 50% of patients within three years (Ottesen et al, 1992; Page et al, 1982). Most of the molecular alterations which have been reported in pre-malignant breast disease have been observed in cases of comedo DCIS (Poller et al, 1993; Radford et al, 1993; and, Tsuda et al, 1993). Non-comedo DCIS is detected by microscopic analysis of breast aspirates or biopsies and is associated with a 10 fold increased risk of breast cancer,

Widespread application of mammography has changed the relative incidence of comedo and non-comedo DCIS such that NCDCIS now represents the predominant form of DCIS diagnosed in the United States (Ottescn et al, 1992; Page et al, 1982; and Pierce et al, 1992). Both forms of DCIS generally recur as invasive cancer at the same site as the pre-malignant lesion (without definitive therapy). The precursor lesions to DCIS are probably atypical ductal hyperplasia and proliferative disease without atypia which are associated with lower rates of breast cancer development, but show further increased risk when associated with a family history of breast cancer (Dupont et al, 1985; Dupont et al, 1989; Dupont et al, 1993; Lawrence, 1990; London et al, 1991; Page et al, 1982; Page et al, 1985; Page et al, 1991; Page et al, 1978; Simpson et al, 1992; Solin et al, 1991; Swain, 1992; Weed et al, 1990).

which corresponds to a 25-30% absolute risk of breast cancer within 15 years (Ottesen

et al, 1992; Page et al, 1982; and, Ward et al, 1992).

What is needed, then, is a sensitive method of detection and diagnosis of breast cancer when the cancerous cells are still in the pre-invasive stage. To illustrate the usefulness in early breast cancer detection of a marker gene and its encoded protein,

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consider the dramatic impact that prostate specific antigen has had on early stage prostate cancer. This method of early detection and diagnosis of breast cancer is presently lacking in the prior art.

Breast cancer occurs in hereditary and sporadic forms. Recently the BRCA 1 gene has been cloned and shown to be mutated in kindreds with hereditary breast and ovarian cancer (Hall et al. 1990, Miki, Y. et al. 1994, Friedman et al. 1994, Castilla et al. 1994, Simard et al. 1994). Although 92% of families with two or more cases of early-onset breast cancer and two cases of ovarian cancer have germ-line mutations in BRCA 1 (Narod et al. in press), the gene has not been shown to be mutated in any truly sporadic case to date (Futreal et al. 1994). Despite the surprising paucity of somatically acquired mutations in sporadic breast cancer, it is still a likely tumor suppressor gene with a key role in breast epithelial cell biology. The BRCA 1 gene encodes a protein of 1863 amino acids with a predicted zinc finger domain observed in proteins which regulate gene transcription. Until the discovery of the function of the BRCA1 gene in conjuction with the delopment of the present invention, the function was unknown.

DISCLOSURE OF THE INVENTION

Epidemiologic studies have established that NCDCIS of the breast is associated with a ten-fold increased risk of breast cancer (absolute risk of 25-30%). It seems likely that this pre-invasive lesion is a determinate precursor of breast cancer because the subsequent development of breast cancer is regularly in the same region of the same breast in which the NCDCIS lesion was found. Important aspects of the present invention concern isolated DNA segments and those isolated DNA segments inserted into recombinant vectors encoding differentially expressed marker genes in abnormal tissue, specifically in NCDCIS, as compared with those expressed in normal tissue, and the creation and use of recombinant host cells through the application of DNA technology, which express these differentially expressed marker genes (Sambrook et al, 1989).

Because there are no cell lines or animal models which clearly display known characteristics of pre-invasive breast disease, human breast tissue samples are essential

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for studying pre-invasive breast disease. Using human tissue samples, we subsequently have developed a method for cDNA cloning from histologically identified lesions in human breast biopsies. We have used this method to clone genes which are differentially expressed in pre-invasive breast lesions such as NCDCIS lesions as compared to genes expressed in normal tissue. The differentially expressed genes detected in pre-invasive breast cancer are called marker genes. Identification of marker genes for pre-invasive breast disease provides improved methods for detection and diagnosis of pre-invasive breast cancer tissue, and further provides marker genes for studies of the molecular events involved in progression from pre-invasive to malignant breast disease.

Analysis of marker gene expression in NCDCIS presents the advantage that cancerous breast tissue at that stage is non-invasive. Detection and diagnosis of NCDCIS by means of differentially expressed marker genes compared to the same marker genes in normal breast tissue, would allow a greater ability to detect, prevent and treat the disease before it becomes invasive and metastasizes. The stage or intermediate condition of NCDCIS is a particularly good candidate for early intervention because it is 1) prior to any invasion and thus prior to any threat to life; 2) it is followed by invasive carcinoma in over 30% of cases if only treated by biopsy; and, 3) there is a long "window" of opportunity (4-8 years) approximately before invasive neoplasia occurs. Thus, NCDCIS is an ideal target for early diagnosis. While these morphologically defined intermediate endpoints have been widely accepted, progress in defining the molecular correlates of these lesions has been hampered by an inability to identify and sample them in a manner which would allow the application of molecular techniques.

Frozen tissue blocks from breast biopsies were used to construct and screen cDNA libraries prepared from NCDCIS tissue, normal breast tissue, breast cancer tissue, and normal human breast epithelial cells. Several cDNAs which were differentially expressed in human DCIS epithelial cells compared to normal breast epithelial cells were cloned and sequenced. One gene which is differentially expressed is the M2 subunit of RibRed which is expressed at low levels in human breast epithelial cells but at higher levels in 4 out of 5 DCIS tissue samples. It is presumed that the

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altered morphologic appearance and determinant biologic behavior of DCIS results from altered expression of genes (such as RibRed) which is important in the induction of breast cancer in humans.

This invention, therefore, provides a method of detecting and diagnosing pre-invasive breast cancer by analyzing marker genes which are differentially expressed in non-comedo DCIS cells. Histopathologic studies have demonstrated that these morphologic patterns in breast tissue lead to invasive breast cancer in at least 20-30% of patients. The present method analyzes gene expression in normal, pre-malignant and malignant breast biopsies; and, it allows simultaneous comparison and cloning of marker genes which are differentially expressed in pre-invasive breast cancer. These marker genes can then be used as probes to develop other diagnostic tests for the early detection of pre-invasive breast cancer.

The present invention concerns DNA segments, isolatable from both normal and abnormal human breast tissue, which are free from total genomic DNA. The isolated DCIS-1 protein product is the regulatory element of the RibRed enzyme. This and all other isolatable DNA segments which are differentially expressed in preinvasive breast cancer can be used in the detection, diagnosis and treatment of breast cancer in its earliest and most easily treatable stages. As used herein, the term "abnormal tissue" refers to pre-invasive and invasive breast cancer tissue, as exemplified by collected samples of non-comedo or comedo DCIS tissues.

As used herein, the term "DNA segment" refers to a DNA molecule which has been isolated free of total genomic DNA of a particular species. Therefore, a DNA segment encoding a differentially expressed protein (as measured by the expression of mRNA) in abnormal tissue refers to a DNA segment which contains differentially expressed-coding sequences in abnormal tissue as compared to those expressed in normal tissue, yet is isolated away from, or purified free from, total genomic DNA of Homo sapiens sapiens. Furthermore, a DNA segment encoding a BRCA1 protein refers to a DNA segment which contains BRCA1 coding sequences, yet is isolated away from, or purified free from, total genomic DNA of Homo sapiens sapiens. Included within the term "DNA segment", are DNA segments and smaller fragments of such segments, and also recombinant vectors, including, for example, plasmids, cosmids,

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phage, viruses, and the like.

Similarly, a DNA segment comprising an isolated or purified differentially expressed gene or comprising an isolated or purified BRCA1 gene refers to a DNA segment including differentially expressed coding sequences or BRCA1 coding sequences isolated substantially away from other naturally occurring genes or protein encoding sequences. In this respect, the term "gene" is used for simplicity to refer to a functional protein, polypeptide or peptide encoding unit. As will be understood by those in the art, this functional term includes both genomic sequences and cDNA sequences. "Isolated substantially away from other coding sequences" means that the gene of interest, in this case, any differentially expressed marker gene or the BRCA1 gene, forms the significant part of the coding region of the DNA segment, and that the DNA segment does not contain large portions of naturally-occurring coding DNA, such as large chromosomal fragments or other functional genes or cDNA coding regions. Of course, this refers to the DNA segment as originally isolated, and does not exclude genes or coding regions later added to the segment by the hand of man.

In particular embodiments, the invention concerns isolated DNA segments and recombinant vectors incorporating DNA sequences which encode differentially expressed genes in pre-invasive breast cancer, each which includes within its amino acid sequence an amino acid sequence in accordance with SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, all seq id no:s 1-7 are derived from non-comedo DCIS samples from Homo sapiens sapiens. In other particular embodiments, the invention concerns isolated DNA segments and recombinant vectors incorporating DNA sequences which encode the M2 subunit of human RibRed that includes within its amino acid sequence the similar amino acid sequence of hamster RibRed corresponding to the M2 subunit of hamster RibRed.

In certain embodiments, the invention concerns isolated DNA segments and recombinant vectors which partially or wholly encode a protein or peptide that includes within its amino acid sequence an amino acid sequence essentially as partially or wholly encoded, respectively, by SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7. Naturally, where the DNA segment or vector encodes a full length differentially expressed protein, or is intended

for use in expressing the differentially expressed protein, the most preferred sequences are those which are essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7 and which encode a protein that exhibits differential expression, e.g., as may be determined by the differential display or differential sequencing assay, as disclosed herein.

The term "a sequence essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7" means that the sequence substantially corresponds to a portion of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, respectively, and has relatively few nucleotides which are not identical to, or a biologically functional equivalent of, the nucleotides of the respective SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7. The term "biologically functional equivalent" is well understood in the art and is further defined in detail herein, for example see pages 24 through 25. Accordingly, sequences which have between about 70% and about 80%; or more preferably, between about 81% and about 90%; or even more preferably, between about 91% and about 99%; of amino acids which are identical or functionally equivalent to the amino acids of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7 will be sequences which are "essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7", respectively.

In particular embodiments, the invention concerns a drug screening method and a gene therapy method that use isolated DNA segments and recombinant vectors incorporating DNA sequences which encode a protein that includes within its amino acid sequence an amino acid sequence in accordance with SEQ ID NO:49, SEQ ID NO:49 derived from breast tissue from Homo sapiens. In other particular embodiments, the invention concerns isolated DNA sequences and recombinant DNA vectors incorporating DNA sequences wich encode a protein taht includes with its amino acid sequence the amino acid sequence of the BRCA1 gene product from human breast tissue.

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In certain embodiments, the invention concerns methods using isolated DNA segments and recombinant vectors which partially or wholly encode a protein or peptide that includes within its amino acid sequence an amino acid sequence essentially as set forth in SEQ ID NO:49. Naturally, where the DNA segment or vector encodes a full length BRCA1 protein, or is intended for use in expressing the BRCA1 protein, the most preferred sequences are those which are essentially as set forth in SEQ ID NO:47 and which encode a protein that retains activity as a negative growth regulator in human breast cells, as may be determined by antisense assay, as disclosed herein.

The term "a sequence essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7" means that the sequence substantially corresponds to a portion of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7, respectively, and has relatively few nucleotides which are not identical to, or a biologically functional equivalent of, the nucleotides of the respective SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7. The term "biologically functional equivalent" is well understood in the art and is further defined in detail herein, for example see pages 24 through 25. Accordingly, sequences which have between about 70% and about 80%; or more preferably, between about 81% and about 90%; or even more preferably, between about 91% and about 99%; of amino acids which are identical or functionally equivalent to the amino acids of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7 will be sequences which are "essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, or SEQ ID NO:7", respectively.

The term "a sequence essentially as set forth in SEQ ID NO:49" means that the sequence substantially corresponds to a portion of SEQ ID NO:49 and has relatively few amino acids which are not identical to, or a biologically functional equivalent of, the nucleotides of SEQ ID NO:49. The term "biologically functional equivalent" is well understood in the art and is further defined in detail herein, for example see pages 24 through 25. Accordingly, sequences which have between about 70% and about 80%; or more preferably, between about 81% and about 90%; or even more

preferably, between about 91% and about 99%; of amino acids which are identical or functionally equivalent to the amino acids of SEQ ID NO:49 will be sequences which are "essentially as set forth in SEQ ID NO:49".

In certain other embodiments, the invention concerns isolated DNA segments and recombinant vectors that include within their sequence a nucleic acid sequence essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7. The term "essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7" is used in the same sense as described above and means that the nucleic acid sequence substantially corresponds to a portion of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7, respectively, and has relatively few codons which are not identical, or functionally equivalent, to the codons of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7, respectively. Again, DNA segments which encode proteins exhibiting differential expression will be most preferred. The term "functionally equivalent codon" is used herein to refer to codons that encode the same amino acid, such as the six codons for arginine or serine, and also refers to codons that encode biologically equivalent amino acids (see Figure 8).

drugs and a gene therapy method which involve the use of isolated DNA segments and recombinant vectors that include within their sequence a nucleic acid sequence essentially as set forth in SEQ ID NO:47 and SEQ ID NO:48. The term "essentially as set forth in SEQ ID NO:47 and SEQ ID NO:48" is used in the same sense as described above and means that the nucleic acid sequence substantially corresponds to a portion of SEQ ID NO:47 and SEQ ID NO:48 respectively, and has relatively few codons which are not identical, or functionally equivalent, to the codons of SEQ ID NO:47 and SEQ ID NO:48, respectively. Again, DNA segments which encode proteins exhibiting the negative regulatory activity of the BRCA1 will be most

preferred. The term "functionally equivalent codon" is used herein to refer to codons

In certain other embodiments, the invention concerns a method for screening

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that encode the same amino acid, such as the six codons for arginine or serine, and also refers to codons that encode biologically equivalent amino acids (see Figure 8).

It will also be understood that amino acid and nucleic acid sequences may include additional residues, such as additional N- or C-terminal amino acids or 5' or 3' sequences, and yet still be essentially as set forth in one of the sequences disclosed herein, so long as the sequence meets the criteria set forth above, including the maintenance of biological protein activity where protein expression is concerned. The addition of terminal sequences particularly applies to nucleic acid sequences which may, for example, include various non-coding sequences flanking either of the 5' or 3' portions of the coding region or may include various internal sequences, i.e., introns, which are known to occur within genes.

Excepting intronic or flanking regions, and allowing for the degeneracy of the genetic code, sequences which have between about 20% and about 50%; or more preferably, between about 50% and about 70%; or even more preferably, between about 70% and about 99%; of nucleotides which are identical to the nucleotides of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7 will be sequences which are "essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7", respectively. Sequences which are essentially the same as those set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7 may also be functionally defined as sequences which are capable of hybridizing to a nucleic acid segment containing the complement of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7, respectively, under relatively stringent conditions. Suitable relatively stringent hybridization conditions will be well known to those of skill in the art (Sambrook et al, 1989).

Excepting intronic or flanking regions, and allowing for the degeneracy of the genetic code, sequences which have between about 20% and about 50%; or more preferably, between about 50% and about 70%; or even more preferably, between about 70% and about 99%; of nucleotides which are identical to the nucleotides of SEQ ID NO:47 and SEQ ID NO:48 will be sequences which are "essentially as set forth in

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SEQ ID NO:47 and SEQ ID NO:48", respectively. Sequences which are essentially the same as those set forth in SEQ ID NO:47 and SEQ ID NO:48 may also be functionally defined as sequences which are capable of hybridizing to a nucleic acid segment containing the complement of SEQ ID NO:47 and SEQ ID NO:48, respectively, under relatively stringent conditions. Suitable relatively stringent hybridization conditions will be well known to those of skill in the art (Sambrook et al, 1989).

It is also important to understand the molecular events which lead to progression from pre-invasive to invasive breast cancer. Breast cancer is a disease that is presumed to involve a series of genetic alterations that confer increasing growth independence and metastatic capability on somatic cells. Identifying the molecular events that lead to the initial development of a neoplasm is therefore critical to understanding the fundamental mechanisms by which tumors arise and to the selection of optimal targets for gene therapy and chemopreventive agents. As intermediate endpoints in neoplastic development, some pre-malignant breast lesions represent important, and possibly rate-limiting steps in the progression of human breast cancer, and careful epidemiological studies have established the relative risk for breast cancer development for specific histologic lesions. In particular, invasive breast cancer develops in the region of the previous biopsy site in at least 25-30% of patients following diagnosis of non-comedo DCIS providing strong evidence that this pre-malignant lesion is a determinant event in breast cancer progression. While these morphologically defined intermediate endpoints have been widely accepted, progress in defining the molecular correlates of these lesions has been hampered by an inability to identify and sample them in a manner which would allow the application of molecular techniques.

The present invention includes a comparison of gene expression between multiple breast tissue biopsy samples as a means to identify differentially expressed genes in pre-malignant breast disease compared with normal breast tissue. These genetic markers should be extremely useful reagents for early diagnosis of breast cancer, and for the delineation of molecular events in progression of breast cancer.

Identification of gene markers which are expressed in the majority of preinvasive breast cancer tissue samples involves cDNA library preparation from both normal and abnormal tissue. This is followed by either a modified differential display method or a differential screening method to identify differential expression of genes which is subsequently confirmed by RT-PCR, nuclease protection assays and in situ hybridization of DCIS tissue RNA and control tissue RNAs (Sambrook et al, 1989). Use of genetic engineering methods can bias the screening to specifically identify genes whose encoded proteins are secreted or are present at the cell surface, in order to find proteins which will be useful markers for diagnostic blood tests (secreted proteins) or for diagnostic imaging studies (cell surface proteins).

Thus, the method of the present invention begins with the collection of at least one tissue sample by a microscopically-directed collection step in which a punch biopsy is obtained exclusively from abnormal tissue which exhibits histological or cytological characteristics of pre-invasive breast cancer. Preferably, the sample site will be an isolatable tissue structure, such as ductal epithelial cells from pre-invasive breast cancer tissue. The mRNA is purified from the sample. Then, a cDNA library is prepared from the mRNA purified from the abnormal tissue sample (Sambrook et al, 1989).

A normal tissue sample is then obtained from the patient, using a sample site. from an area of tissue which does not exhibit histological or cytological characteristics of pre-invasive cancer. A cDNA library is also prepared from this normal tissue sample.

The abnormal tissue cDNA library can then be compared with the normal tissue cDNA library by differential display or differential screening to determine whether the expression of at least one marker gene in the abnormal tissue sample is different from the expression of the same marker gene in the normal tissue sample.

Further diagnostic steps can be added to the method by cloning the marker gene using sequence-based amplification to create a cloned marker gene which can then be DNA-sequenced in order to derive the protein sequence. The protein sequence is then used to generate antibodies which will recognize these proteins by antibody recognition of the antigen. The presence of the antibody-recognized antigen can then be detected by means of conventional medical diagnostic tests.

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This invention also includes methods of screening for compounds and gene therapy methods using the BRCA1 gene. BRCA1 mRNA is expressed at 5-10 fold higher levels in normal mammary tissue than in invasive breast cancer samples. Having demonstrated that mRNA expression levels of BRCA1 are higher in normal mammary cells than in cancer cells, antisense methods were used to test the hypothesis that BRCA1 expression inhibits cell growth. These tests showed that diminished expression of BRCA1 increased the proliferative rate of breast cells.

An object of the present invention, then, is to provide a method of early detection of pre-invasive breast cancer in human tissue.

It is a further object of this invention to identify early marker genes for preinvasive breast disease which can be used in screening methods for early pre-invasive breast cancer.

It is also an object of this invention to produce a cDNA library from preinvasive breast cancer tissue resulting in a permanent genetic sample of that preinvasive breast cancer tissue.

It is also an object of this invention to provide a drug or biological screening method using the BRCA 1 promoter region and gene therapy method using the BRCA 1 gene.

		List of Abbreviations		
20	TPA	Phorbol 12-myristate 13-acetate		
	MCF-7	An immortalized cell line derived from a metastasis of		
		human breast cancer		
	HMEC	A primary (non-immortalized) cell line derived from		
		breast epithelial cells obtained during reduction		
25		mammoplasty ,		
	DCIS	Ductal Carcinoma-in-situ		
	NCDC	Non-Comedo Ductal Carcinoma in situ		
	cDNA	Complementary DNA obtained from an RNA template		
	DNA	Deoxyribonucleic Acid		
30 RT-PCR		Reverse Transcriptase-Polymerase Chain Reaction		
	RibRed	Ribonucleotide Reductase		

- Fig. 1 shows Table I which describes anatomic lesion types in the human breast with pre-malignant implication.
- Fig. 2 shows a model for pre-malignant conditions, highlighting magnitude of risk for progression to clinical malignancy.

Fig. 3 contains color photos of DCIS tissue, before (upper left panel) and after microscopically-directed excisional punch biopsy (upper right panel). The lower panels show tissue samples of normal breast tissue (lower left panel), and invasive breast cancer (lower right panel).

Fig. 4 shows expression of collagen III mRNA in tissue mRNA samples, analyzed by RNase protection assay methods.

Fig. 5 shows differential display of cDNAs obtained from patient tissue samples and controls.

Fig. 6 shows a comparison of the sequence between DCIS-1 and the human and hamster genes.

Fig. 7 shows expression of DCIS-1 mRNA in tissue mRNA samples analyzed by RNase protection assay as described in the legend to Figure 4.

Fig. 8 is Table II which displays the genetic code.

Fig. 9 is a Table which lists differentially expressed marker genes.

Figs. 10A and 10B shows expression of BRCA1 mRNA during breast cancer progression by PCR detection and nuclease protection assay, respectively.

Figs. 11A and 11B is a comparison of BRCA1 expression in normal breast and invasive breast cancer using nuclease protection assay of RNA, respectively.

Figs. 12A, 12B, and 12C show that antisense inhibition of BRCA1 accelerates mammary cell proliferation.

Figs. 13A and 13B includes a Northern blot of mRNA and nuclear runon studies that show that ribonucleotide reductase M2 mRNA is cell cycle regulated in MCF-7 cells.

Fig. 14 includes a nuclease protection assay that shows that antisense inhibition of BRCA1 in human mammary cells decreases BRCA1 mRNA and increases ribonucleotide reductase mRNA.

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UTILITY STATEMENT

The detection of differentially expressed genes in pre-invasive breast tissue, specifically in non-comedo ductal carcinoma in situ as compared to genes expressed in normal tissue, is useful in the diagnosis, prognosis and treatment of human breast cancer. Such differentially expressed genes are effective marker genes indicating the significantly increased risk of breast cancer in a patient expressing these differentially expressed marker genes. These marker genes are useful in the detection, early diagnosis, and treatment of breast cancer in humans.

The discovery of the function of the BRCA 1 gene has broad utility including, in the present invention, development of methods to treat familial and sporadic breast cancers as well as screen for therapeutic drugs through production of important indicator compounds.

ACTIVITY STATEMENT

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Of the differentially expressed genes described in this invention, DCIS-1 encodes a gene similar to the M2 subunit of hamster ribonucleotide reductase. The M2 subunit of ribonucleotide reductase (RibRed, hereafter) is responsible for regulation of RibRed. The differential levels of expression of the marker genes described in this invention (Seq ID No.s 1-7), indicate genetic changes which have been linked to the presence of pre-invasive breast cancer.

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The BRCA1 gene (Seq. ID No. 47) is differentially expressed in invasive breast cancer cells. The BRCA1 gene product is a negative regulator of mammary cell proliferation which is expressed at diminished levels in sporadic breast cancer.

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BEST MODE FOR CARRYING OUT THE INVENTION

For the purposes of the subsequent description, the following definitions will be used:

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Nucleic acid sequences which are "complementary" are those which are capable of base-pairing according to the standard Watson-Crick complementarity rules. That is, that the larger purines will always base pair with the smaller pyrimidines to form only combinations of Guanine paired with Cytosine (G:C) and Adenine paired with

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either Thymine (A:T) in the case of DNA or Adenine paired with Uracil (A:U) in the case of RNA.

"Hybridization techniques" refer to molecular biological techniques which involve the binding or hybridization of a probe to complementary sequences in a polynucleotide. Included among these techniques are northern blot analysis, southern blot analysis, nuclease protection assay, etc.

"Hybridization" and "binding" in the context of probes and denatured DNA are used interchangeably. Probes which are hybridized or bound to denatured DNA are aggregated to complementary sequences in the polynucleotide. Whether or not a particular probe remains aggregated with the polynucleotide depends on the degree of complementarity, the length of the probe, and the stringency of the binding conditions. The higher the stringency, the higher must be the degree of complementarity and/or the longer the probe.

"Probe" refers to an oligonucleotide or short fragment of DNA designed to be sufficiently complementary to a sequence in a denatured nucleic acid to be probed and to be bound under selected stringency conditions.

"Label" refers to a modification to the probe nucleic acid that enables the experimenter to identify the labeled nucleic acid in the presence of unlabeled nucleic acid. Most commonly, this is the replacement of one or more atoms with radioactive isotopes. However, other labels include covalently attached chromophores, fluorescent moeities, enzymes, antigens, groups with specific reactivity, chemiluminescent moeities, and electrochemically detectable moeities, etc.

"Marker gene" refers to any gene selected for detection which displays differential expression in abnormal tissue as opposed to normal tissue. It is also referred to as a differentially expressed gene.

"Marker protein" refers to any protein encoded by a "marker gene" which protein displays differential expression in abnormal tissue as opposed to normal tissue.

"Tissuemizer" describes a tissue homogenization probe.

"Abnormal tissue" refers to pathologic tissue which displays cytologic, histologic and other defining and derivative features which differ from that of normal

tissue. This includes in the case of abnormal breast tissue, among others, pre-invasive and invasive neoplasms.

"Normal tissue" refers to tissue which does not display any pathologic traits.

"PCR technique" describes a method of gene amplification which involves sequenced-based hybridization of primers to specific genes within a DNA sample (or library) and subsequent amplification involving multiple rounds of annealing, elongation and denaturation using a heat-stable DNA polymerase.

"RT-PCR" is an abbreviation for reverse transcriptase-polymerase chain reaction. Subjecting mRNA to the reverse transcriptase enzyme results in the production of cDNA which is complementary to the base sequences of the mRNA. Large amounts of selected cDNA can then be produced by means of the polymerase chain reaction which relies on the action of heat-stable DNA polymerase produced by Thermus aquaticus for its amplification action.

"Microscopically-directed" refers to the method of tissue sampling by which the tissue sampled is viewed under a microscope during the sampling of that tissue such that the sampling is precisely limited to a given tissue type, as the investigator requires. Specifically, it is a collection step which involves the use of a punch biopsy instrument. This surgical instrument is stereotactically manually-directed to harvest exclusively from abnormal tissue which exhibits histologic or cytologic characteristics of pre-invasive cancer. The harvest is correlated with a companion slide, stained to recognize the target tissue.

"Differential display" describes a method in which expressed genes are compared between samples using low stringency PCR with random oligonucleotide primers.

"Differential screening" describes a method in which genes within cDNA libraries are compared between two samples by differential hybridization of cDNAs to probes prepared from each library.

"Nuclease protection assay" refers to a method of RNA quantitation which employs strand specific nucleases to identify specific RNAs by detection of duplexes.

"Differential expression" describes the phenomenon of differential genetic expression seen in abnormal tissue in comparison to that seen in normal tissue.

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"Isolatable tissue structure" refers to a tissue structure which when visualized microscopically or otherwise is able to be isolated from other different surrounding tissue types.

"In situ hybridization of RNA" refers to the use of labeled DNA probes employed in conjunction with histological sections on which RNA is present and with which the labeled probe can hybridize allowing an investigator to visualize the location of the specific RNA within the cell.

"Comedo DCIS cells" refers to cells comprising an in situ lesion with the combined features of highest grade DCIS.

"Non-comedo DCIS cells" refers to cells of DCIS lesions without comedo features.

"Cloning" describes separation and isolation of single genes.

"Sequencing" describes the determination of the specific order of nucleic acids in a gene or polynucleotide.

The present invention provides a method for detecting and diagnosing cancer by analyzing marker genes which are differentially expressed in early, pre-invasive breast cancer, specifically in non-comedo DCIS cells. Our histopathologic studies have demonstrated that certain morphologic patterns in breast tissue are pre-malignant; leading to invasive breast cancer in at least 20-30% of patients. We have developed a new method for analyzing gene expression in normal, pre-malignant and malignant breast biopsies which allows simultaneous comparison and cloning of marker genes which are differentially expressed in pre-invasive breast cancer. These marker genes (which appear as differentially expressed genes in pre-invasive breast cancer) can be used as probes to develop diagnostic tests for the early detection of pre-invasive breast cancer (Sambrook, 1989).

The present invention thus comprises a method of identification of marker genes which are expressed in the majority of pre-invasive breast cancer tissue samples. It involves cDNA library preparation followed by a modified differential display method. Use of genetic engineering methods (Sambrook, 1989) can bias the screening to specifically identify genes whose encoded proteins are secreted or are present at the cell

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surface, in order to find proteins which will be useful markers for diagnostic blood tests (secreted proteins) or for diagnostic imaging studies (cell surface proteins).

Naturally, the present invention also encompasses DNA segments which are complementary, or essentially complementary, to the sequence set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:47 and SEQ ID NO:48. Nucleic acid sequences which are "complementary" are those which are capable of base-pairing according to the standard Watson-Crick complementarity rules. As used herein, the term "complementary sequences" means nucleic acid sequences which are substantially complementary, as may be assessed by the same nucleotide comparison set forth above, or as defined as being capable of hybridizing to the nucleic acid segment of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:47 and SEQ ID NO:48 under relatively stringent conditions such as those described herein.

The nucleic acid segments of the present invention, regardless of the length of the coding sequence itself, may be combined with other DNA sequences, such as promoters, polyadenylation signals, additional restriction enzyme sites, multiple cloning sites, other coding segments, and the like, such that their overall length may vary considerably. It is therefore contemplated that a nucleic acid fragment of almost any length may be employed, with the total length preferably being limited by the ease of preparation and use in the intended recombinant DNA protocol. For example, nucleic acid fragments may be prepared which include a short stretch complementary to SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:47 and SEQ ID NO:48, such as about 10 nucleotides, and which are up to 10,000 or 5,000 base pairs in length, with segments of 500 being preferred in most cases. DNA segments with total lengths of about 1,000, 500, 200, 100 and about 50 base pairs in length are also contemplated to be useful.

It will also be understood that this invention is not limited to the particular nucleic acid and amino acid sequences of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:47, SEQ ID NO:48, and SEQ ID NO:49. Recombinant vectors and isolated DNA

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segments may therefore variously include the differentially expressed coding regions or the BRCAl coding regions themselves, coding regions bearing selected alterations or modifications in the basic coding region, or they may encode larger polypeptides which nevertheless include differentially expressed-coding regions and the BRCAl coding regions or may encode biologically functional equivalent proteins or peptides which have variant amino acids sequences.

The DNA segments of the present invention encompass biologically functional equivalent differentially expressed proteins and peptides biologically functional equivalent proteins of BRCA1. Such sequences may arise as a consequence of codon redundancy and functional equivalency which are known to occur naturally within nucleic acid sequences and the proteins thus encoded. Alternatively, functionally equivalent proteins or peptides may be created via the application of recombinant DNA technology, in which changes in the protein structure may be engineered, based on considerations of the properties of the amino acids being exchanged. Changes designed by man may be introduced through the application of site-directed mutagenesis techniques, e.g., to introduce improvements to the antigenicity of the protein or to test site-directed mutants or others in order to examine carcinogenic activity of the differentially expressed marker genes at the molecular level.

If desired, one may also prepare fusion proteins and peptides, e.g., where the differentially expressed marker gene coding regions are aligned within the same expression unit with other proteins or peptides having desired functions, such as for purification or immunodetection purposes (e.g., proteins which may be purified by affinity chromatography and enzyme label coding regions, respectively).

Recombinant vectors form important further aspects of the present invention. Particularly useful vectors are contemplated to be those vectors in which the coding portion of the DNA segment is positioned under the control of a promoter. The promoter may be in the form of the promoter which is naturally associated with a RIBRED gene, e.g., in human cells, as may be obtained by isolating the 5' non-coding sequences located upstream of the coding segment or exon, for example, using recombinant cloning and/or PCR technology, in connection with the compositions disclosed herein.

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In other embodiments, it is contemplated that certain advantages will be gained by positioning the coding DNA segment under the control of a recombinant, or heterologous, promoter. As used herein, a recombinant or heterologous promoter is intended to refer to a promoter that is not normally associated with a differentially expressed marker gene or the BRCA1 gene in its natural environment. Such promoters may include MMTV promoters normally associated with other genes, and/or promoters isolated from any other bacterial, viral, eukaryotic, or mammalian cell. Naturally, it will be important to employ a promoter that effectively directs the expression of the DNA segment in the cell type chosen for expression. The use of promoter and cell type combinations for protein expression is generally known to those of skill in the art of molecular biology, for example, see Sambrook et al. (1989). The promoters employed may be constitutive, or inducible, and can be used under the appropriate conditions to direct high level expression of the introduced DNA segment, such as is advantageous in the large-scale production of recombinant proteins or peptides. Appropriate promoter systems contemplated for use in high-level expression include, but are not limited to appropriate bacterial promoters.

As mentioned above, in connection with expression embodiments to prepare recombinant differentially expressed marker gene encoded proteins and peptides, it is contemplated that longer DNA segments will most often be used, with DNA segments encoding the entire differentially expressed protein or subunit being most preferred. However, it will be appreciated that the use of shorter DNA segments to direct the expression of differentially expressed peptides or epitopic core regions, such as may be used to generate anti-marker protein antibodies, also falls within the scope of the invention (Harlow et al, 1988).

amino acids in length, or more preferably, from about 15 to about 30 amino acids in length are contemplated to be particularly useful. The C terminus of proteins provide an excellent region for peptide antigen recogition (Harlow et al, 1988). DNA segments encoding peptides will generally have a minimum coding length in the order of about 45 to about 147, or to about 90 nucleotides. DNA segments encoding partial length

DNA segments which encode peptide antigens from about 15 to about 50

peptides may have a minimum coding length in the order of about 50 nucleotides for

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a polypeptide in accordance with seq id no:3, or about 264 nucleotides for a polypeptide in accordance with SEQ ID NO: 1.

In addition to their use in directing the expression of the differentially expressed marker proteins, the nucleic acid sequences disclosed herein also have a variety of other uses. For example, they also have utility as probes or primers in nucleic acid hybridization embodiments. As such, it is contemplated that oligonucleotide fragments corresponding to the sequences of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7 for stretches of between about 10 to 15 nucleotides and about 20 to 30 nucleotides will find particular utility. Longer complementary sequences, e.g., those of about 40, 50, 100, 200, 500, 1000, and even up to full length sequences of about 2,000 nucleotides in length, will also be of use in certain embodiments.

The ability of such nucleic acid probes to specifically hybridize to differentially expressed marker gene sequences will enable them to be of use in detecting the presence of complementary sequences in a given sample. However, other uses are envisioned, including the use of the sequence information for the preparation of mutant species primers, or primers for use in preparing other genetic constructions.

Nucleic acid molecules having stretches of 20, 30, 50, or even of 500 nucleotides or so, complementary to SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7 are particularly contemplated as hybridization probes for use in, e.g., Southern and Northern blotting. This would allow differentially expressed structural or regulatory genes to be analyzed, both in patients and sample tissue from pre-invasive and invasive breast tissue. The total size of fragment, as well as the size of the complementary stretch(es), will ultimately depend on the intended use or application of the particular nucleic acid segment. Smaller fragments will generally find use in hybridization embodiments, wherein the length of the complementary region may be varied, such as between about 10 and about 100 nucleotides, but larger complementary stretches of up to about 300 nucleotides may be used, according to the length complementary sequences one wishes to detect.

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Nucleic Acid Hybridization

The use of a hybridization probe of about 10 nucleotides in length allows the formation of a duplex molecule that is both stable and selective. Molecules having complementary sequences over stretches greater than 10 bases in length are generally preferred, though, in order to increase stability and selectivity of the hybrid, and thereby improve the quality and degree of specific hybrid molecules obtained. One will generally prefer to design nucleic acid molecules having gene-complementary stretches of 15 to 20 nucleotides, or even longer where desired.

Hybridization probes may be selected from any portion of any of the sequences

disclosed herein. All that is required is to review the sequences set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7 and to select any continuous portion of one of the sequences, from about 10 nucleotides in length up to and including the full length sequence, that one wishes to utilise as a probe or primer. The choice of probe and primer sequences may be governed by various factors, such as, by way of example only, one may wish to employ primers from towards the termini of the total sequence, or from the ends of the functional domain-encoding sequences, in order to amplify further DNA; one may employ probes corresponding to the entire DNA, or to the 5' region, to clone marker-type genes from other species or to clone further marker-like or homologous genes from any species including human; and one may employ randomly selected, wild-type and mutant probes or primers with sequences centered around the RibRed M2 subunit encoding sequence to screen DNA samples for differentially expressed levels of

The process of selecting and preparing a nucleic acid segment which includes a sequence from within SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7 may alternatively be described as "preparing a nucleic acid fragment". Of course, fragments may also be obtained by other techniques such as, e.g., by mechanical shearing or by restriction enzyme digestion. Small nucleic acid segments or fragments may be readily prepared by, for example, directly synthesizing the fragment by chemical means, as is commonly

RibRed, such as to identify human subjects which may be expressing differential levels

of RibRed and thus may be susceptible to breast cancer.

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practiced using an automated oligonucleotide synthesizer. Also, fragments may be obtained by application of nucleic acid reproduction technology, such as the PCR technology of U.S. Patent 4,603,102 (incorporated herein by reference), by introducing selected sequences into recombinant vectors for recombinant production, and by other recombinant DNA techniques generally known to those of skill in the art of molecular biology.

Accordingly, the nucleotide sequences of the invention may be used for their ability to selectively form duplex molecules with complementary stretches of differentially expressed marker genes or cDNAs. Depending on the application envisioned, one will desire to employ varying conditions of hybridization to achieve varying degrees of selectivity of probe towards target sequence. For applications requiring high selectivity, one will typically desire to employ relatively stringent conditions to form the hybrids, e.g., one will select relatively low salt and/or high temperature conditions, such as provided by 0.02M-0.15M NaCl at temperatures of 50°C to 70°C. Such selective conditions tolerate little, if any, mismatch between the probe and the template or target strand, and would be particularly suitable for isolating specific differentially expressed marker genes.

Of course, for some applications, for example, where one desires to prepare mutants employing a mutant primer strand hybridized to an underlying template or where one seeks to isolate marker gene sequences from related species, functional equivalents, or the like, less stringent hybridization conditions will typically be needed in order to allow formation of the heteroduplex. In these circumstances, one may desire to employ conditions such as 0.15M-0.9M salt, at temperatures ranging from 20°C to 55°C. Cross-hybridizing species can thereby be readily identified as positively hybridizing signals with respect to control hybridizations. In any case, it is generally appreciated that conditions can be rendered more stringent by the addition of increasing amounts of formamide, which serves to destabilize the hybrid duplex in the same manner as increased temperature. Thus, hybridization conditions can be readily manipulated, and thus will generally be a method of choice depending on the desired results.

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In certain embodiments, it will be advantageous to employ nucleic acid sequences of the present invention in combination with an appropriate means, such as a label, for determining hybridization. A wide variety of appropriate indicator means are known in the art, including fluorescent, radioactive, enzymatic or other ligands, such as avidin/biotin, which are capable of giving a detectable signal. In preferred embodiments, one will likely desire to employ a fluorescent label or an enzyme tag, such as urease, alkaline phosphatase or peroxidase, instead of radioactive or other environmental undesirable reagents. In the case of enzyme tags, colorimetric indicator substrates are known which can be employed to provide a means visible to the human eye or spectrophotometrically, to identify specific hybridization with complementary nucleic acid-containing samples.

In general, it is envisioned that the hybridization probes described herein will be useful both as reagents in solution hybridization as well as in embodiments employing a solid phase. In embodiments involving a solid phase, the test DNA (or RNA) is adsorbed or otherwise affixed to a selected matrix or surface. This fixed, single-stranded nucleic acid is then subjected to specific hybridization with selected probes under desired conditions. The selected conditions will depend on the particular circumstances based on the particular criteria required (depending, for example, on the G+C contents, type of target nucleic acid, source of nucleic acid, size of hybridization probe, etc.). Following washing of the hybridized surface so as to remove nonspecifically bound probe molecules, specific hybridization is detected, or even quantified, by means of the label. (Sambrook et al., 1989).

In a preferred embodiment of the method, certain preliminary procedures are necessary to prepare the sample tissue and the probes before the detection of differential expression of marker genes in abnormal tissue as compared to that in normal tissue can be accomplished.

SAMPLE PREPARATION

RNA purification

RNA was isolated from frozen tissue samples by mincing of microdisected frozen tissue fragments with a razor blade and then adding 800 microliter of 5.6M

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guanidinium to increase mixing, followed by a 30 second microcentrifuge centrifugation at 14,000 rpm to remove particulate matter. The supernatant was then removed and the viscosity was reduced by multiple aspirations through a 22 gauge needle and then 200 ul of chloroform was added and the sample was incubated on ice for 15 minutes (during this time the sample was vortexed multiple times). Following incubation with chloroform, the sample was centrifuged for 15 minutes at 14,000 rpm and the aqueous layer was removed and ethanol precipitated. This extraction method produces RNA which is primarily derived from cells of epithelial origin. In order to obtain RNA samples which presumably includes RNA derived from these stromal cells; the particulate material (remaining in the pellet from the 30 second centrifugation) was homogenized with a tissuemizer, washed with PBS, treated with collagenase at 37°C for 30 minutes, sonicated, extracted with phenol/chloroform and ethanol precipitated.

cDNA libraries were constructed in lambda phage using polyA-selected mRNA from the following samples; cultured human breast epithelial cells, tissue from three reduction mammoplasty patients, tissue from three DCIS patients, and tissue from one DCIS patient (patient #10) that showed a focus of microinvasion adjacent to an area of DCIS. Multiple punches were needed to obtain sufficient RNA for polyA selection and library construction. 200 ug of total RNA was obtained by pooling 20 punches from normal breast tissue (reduction mammoplasty samples) and 5-8 punches from DCIS lesions, presumably reflecting the greater cellularity of the DCIS samples. cDNA libraries were constructed by first and second strand cDNA synthesis followed by the addition of directional synthetic linkers (ZAP-cDNA Synthesis Kit, Stratagene, La Jolla, California). The Xho I-Eco Rl linkered cDNA was then ligated into lambda arms, packaged with packaging extracts, and then used to infect XL1-blue bacteria resulting in cDNA libraries.

PROBE PREPARATION

The collagen III probe employed for nuclease protection assays was constructed by subcloning the 208 bp Hinc II-Pst I fragment from the 3' untranslated region of the human type III procollagen gene into pGem4Z. This region of the human procollagen III gene was obtained by PCR amplification of published sequence (Ala-Kokko et al, 1989) followed by restriction with Hinc II and Pst I. For a control probe to assure

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equal loading and recovery of RNA, we used a T7 polymerase-generated probe for human glyceraldehyde phosphate dehydrogenase (GADP) which protects a 140 bp Sac I-Xba I fragment; (a generous gift from Janice Nigro, Vanderbilt University). Probe DCIS-1 was generated by linearizing the rescued plasmid with Pvu II, which should generate a 200 bp protected fragment. RNase protection assays were performed with 1 ug of unselected RNA and the above-cited probes using the methods we have reported previously (Holt, 1993).

Differential Display-based cloning of cDNAs:

Rescued cDNA library samples were used as templates for low stringency PCR with the either a pair of 25 bp primers or an anchored 14 bp primer paired with a random 25 bp primer. Random 25 bp primers were generated by a computer-based algorithm (Jotte and Holt, unpublished). Samples were denatured for two minutes at 95°C followed by 40 cycles, each cycle consisting of denaturation for 1 minute at 94°C., annealing for 2 minutes at 25°C., and extension for 1 minute at 72°C. The samples were then run on an 6% non-denaturing polyacrylamide gel, which was dried and autoradiographed. Specific bands were excised then reamplified with the same primers used for their generation. Specificity was confirmed on 6% polyacrylamide gel, and samples were purified by ethanol precipitation of the remainder of the PCR reaction. Fragments were then individually cloned into Srfl cut vectors by standard methods using PCR-ScriptTMSK(+) Cloning Kit (Stratagene, LaJolla, California) and then sequenced.

EXAMPLE 1

Studies showing Increased Risk of Breast Cancer in Patients with DCIS

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Since the 1970's, studies of pre-invasive lesions associated with the development of breast cancer have been undertaken in an attempt to refine histologic and cytologic criteria for the hyperplastic lesions analogous to those of the uterine cervix and colon. Because of the availability of tissue from breast biopsies done many years previously, cohorts of women who underwent breast biopsies 15 to 20 years ago, can be studied to determine the risk for development of breast cancer attributable to specific lesions.

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Many concurrent studies evaluating lesions associated with cancer at time of cancer diagnosis led the way in pointing out lesions of potential interest (Wellings et al, 1975). Hopefully, these intermediate stages in cancer development will serve to provide indicators of breast cancer development sufficiently precise to guide prevention and intervention strategies (Weed et al, 1990; Lippman et al, 1990). Such intermediate elements prior to the development of metastatic capable cancers also provide the opportunity to define the molecular biology of these elements. Studies of the development of pre-invasive breast disease have provided insight into different types of lesions with different implications for breast cancer risk and the process of carcinogenesis (See Figure 1). Pre-invasive breast disease is herewith defined to be any reproducibly defined condition which confers an elevated risk of breast cancer approaching double that of the general population (Komitowski et al, 1990). The specifically-defined atypical hyperplasias and lobular carcinoma in situ confer relative risks of four to ten times that of the general population. This risk is for carcinoma to develop anywhere in either breast (Page et al, 1985; Page et al, 1991). The statistical significance of these observations have regularly been < .0001. Thus, absolute risk figures of 10-20% likelihood of developing into invasive carcinoma in 10 to 15 years arise. DCIS is a very special element in this story because the magnitude of risk is as high as any other condition noted (P < .00005), but remarkably, the developing invasive cancer is in the same site in the same breast. This local recurrence and evolution to invasiveness marks these lesions as determinate precursors of invasive breast cancer (Betsill et al, 1978; Page et al, 1982). These figures are for the type of DCIS which has become detected very commonly since the advent of mammography, the small and NCDCIS variety. It is likely that the comedo DCIS variety indicates a much greater risk, often presenting as larger lesions, and treated regularly by mastectomy in the past 50 years making follow-up studies impossible (Figure 1).

The precision of histopathologic diagnosis in this area as noted in Table I (shown in Figure 1) was most convincingly confirmed in a large, prospective study (London et al, 1991). There has also been a recent review of the reproducibility of the assignment of diagnosis by a panel of pathologists (Schnitt et al, 1992). The precision has been fostered by combining histologic pattern criteria with cytologic and extent of

lesion criteria. Classic surgical pathology criteria were predominantly derived from histologic pattern only. A further point of relevance to the importance of these histopathologically defined lesions of pre-malignancy in the breast is the relationship to familiality. A family history of breast cancer in a first degree relatives confers about a doubling of breast cancer risk. However, women with the atypical hyperplasias at biopsy and a family history of breast cancer are at 9-10 times the risk of developing invasive breast cancer as the general population (Dupont et al, 1985; Dupont et al, 1989).

Careful consideration of all of the above-mentioned epidemiologic data has led to the following model for progression from generalized pre-malignant lesions to determinant lesions to invasive cancer. Figure 2 shows this model for the induction and progression of pre-invasive breast disease based on study of the Vanderbilt cohort (Dupont et al, 1985) of more than 10,000 breast biopsies (follow-up rate 85%; median

time of 17 years; 135 women developed breast cancer).

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EXAMPLE 2

Identification of genes which are differentially expressed in DCIS Construction of cDNA libraries from DCIS lesions

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In order to study differential gene expression in DCIS, we collected cases of NCDCIS. The diagnosis of DCIS is made on histomorphologic grounds based on architectural, cytologic, and occasionally extent criteria. NCDCIS lacks comedo features and consists of microscopic intraductal lesions which fill and extend the duct, contain rigid internal architecture, and often have hyperchromatic and monomorphic nuclei.

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Study of non-comedo DCIS for differential marker gene expression indicates the diagnostic utility of comparison of marker gene expression in these tissues. Although the morbidity and mortality of breast cancer clearly results from invasion and metastasis, the development of breast cancer is clearly significant in its early stages for two basic reasons:

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The molecular changes will presumably be simpler in early lesions than in later lesions which may have acquired numerous mutations or "hits"; and

 Successful prevention strategies may require attacking cancer before it develops the capacity to invade or metastasize.

Non-comedo DCIS is the earliest determinant lesion which recurs locally as invasive cancer. Although comedo DCIS may be technically easier to study because the tumors are larger, its aggressiveness and the presence of numerous genetic alterations (such as p53 and erbB2) suggest that it may have advanced beyond the earliest stages of carcinogenesis.

The commercial utility of a method for prevention of cancer is clear. In order to study differential gene expression in DCIS, breast tissue with extensive microscopic non-comedo DCIS was identified and banked in a frozen state. cDNA libraries were constructed from mRNA isolated from frozen sections of DCIS lesions. Tissue samples from patients with mammographic results consistent with DCIS were cryostat frozen and a definitive diagnosis was made by the histopathologic criteria which we have described (Jensen et al, Submitted for publication; Holt et al, In press).

Control mRNA was obtained from frozen tissue samples obtained from reduction mammoplasties and from cultured human breast epithelial cells. Because non-comedo DCIS is a microscopic lesion, we had to microlocalize regions of DCIS in biopsy samples. To accomplish this we prepared frozen sections in which we located regions of DCIS and then employed a 2 mm punch to obtain an abnormal tissue sample only from those regions that contained DCIS. This selective harvesting was accomplished by carefully aligning the frozen section slide with the frozen tissue block and identifying areas of interest. The harvest of the appropriate area was then confirmed with a repeat frozen section. A similar approach was used to isolate mRNA from lobules of normal breast in samples collected from a reduction mammoplasty. Prior studies have shown that breast lobules are approximately 2.5 mm in diameter, thus the 2 mm punch provided a well-tailored excision. This microlocation and collection step, in which abnormal tissue samples are collected from an isolatable tissue structure, was performed with extreme care and was absolutely crucial to the success of these studies. Contamination by normal breast epithelial cells or by breast stromal cells would clearly negatively skew the differential screening approach. If the punch biopsy did not cleanly

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excise DCIS without contamination by other cell types or tissues then the sample was not used for mRNA isolation (Jensen et al, Submitted for publication). Figure 3 contains color photos of DCIS (abnormal) tissue, before (upper left panel) and after excisional punch biopsy (upper right panel). The lower panels show tissue samples of normal breast tissue (lower left panel), and invasive breast cancer (lower right panel).

Following microlocation punch harvesting of the frozen tissue, RNA was isolated, purified, and employed to construct cDNA libraries. RNA was isolated following mincing of tissue in 5.6M guanidinium isothiocyanate and 40% phenol, centrifugation to remove particulate matter, viscosity reduction by repeated aspiration through a 22 gauge needle, chloroform extraction and ethanol precipitation. In most samples there was particulate matter resistant to guanidinium-phenol extraction that was white in color and fibrous in appearance and was presumed to represent breast stroma. This stromal material was sparse in DCIS samples but abundant in samples obtained from normal breast tissue derived from reduction mammoplasties. The stromal material was minced with a tissuemizer, washed with PBS, treated with collagenase at 37°C for 30 minutes, sonicated, extracted with phenol/chloroform and ethanol precipitated. 200 ug of total RNA was obtained by pooling 20 punches from normal breast tissue (reduction mammoplasty samples) and 5-8 punches from DCIS lesions, presumably reflecting the greater cellularity of the DCIS samples. All libraries had greater than 50% inserts and contained between 2 X 106 and 7 X 107 phage recombinants with an average insert size varying between 500 and 1000 base pairs.

EXAMPLE 3

Development of an extraction method which produces breast epithelial RNA

It was necessary that tissue samples not be contaminated by non-epithelial stromal cells. Such contamination would complicate efforts to compare gene expression between samples. In order to test the extent of stromal contamination of the mRNA samples, we determined the level of expression of collagen III mRNA by an RNase protection assay. RNase protection assays were employed in these and subsequent studies because it is a quantitative method and can be performed on small amounts of unselected RNA. Collagen III mRNA was identified in the presumed stromal fraction

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of the normal breast tissue and to a lesser extent in the microinvasive breast cancer sample, but no expression of collagen III was detected in the DCIS samples which were subsequently employed for cDNA library construction. Figure 4 compares expression in NL 2 and #10CA with other patient samples and NL1 to determine collagen III expression.

Expression of Collagen III mRNA in tissue mRNA samples was analyzed by RNase protection assay by methods we have reported previously (Holt, 1993). One µg of mRNA was hybridized with two labeled RNA probes: a T7 polymerase-generated probe for human glyceraldehyde phosphate dehydrogenase (GADP) which protects a 140 bp Sac I-Xba I fragment; and a T7 polymerase-generated probe which protects a 208 bp Hinc II-Pst I fragment from the 3' untranslated region of the human type III procollagen gene (Coll III) obtained by PCR subcloning of the published sequence (Ala-Kokko et al, 1991). RNA samples were labeled as follows: NL1 is RNA from cultured human breast epithelial cells (Hammond et al, 1984), NL2 is RNA from normal breast tissue, NL3 is RNA derived from the fibrous stromal fraction of breast tissue as described (Jensen et al, Submitted for publication), NL4 is another sample from normal breast tissue. This is described in greater detail on page 30 of this patent application. #12,#8,#4,#6, and #10 are from patient samples with DCIS. Sample #10CA is RNA obtained from the small focus of microinvasion shown in Figure 3. Con is a control sample using tRNA.

EXAMPLE 4

Screening of cDNA libraries

Following successful testing which demonstrated that stromal contamination was not a problem, cDNA libraries were constructed in lambda phage using polyA-selected mRNA from the following samples: cultured human breast epithelial cells, tissue from three reduction mammoplasty patients, tissue from three DCIS patients, and tissue from one DCIS patient (patient #10) that showed a small focus of invasion adjacent to an area of DCIS. Multiple punches were needed to obtain sufficient RNA for polyA selection and library construction. Selective handling of tissue was accomplished.

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Comparison of gene expression between samples was performed by either differential screening or a modification of differential display (Liang et al, 1992a; Liang et al, 1992b; Saiki et al, 1988; Melton et al, 1984). Plasmid DNA was prepared from the cDNA libraries following helper phage rescue and screened by two independent methods. Figure 5 below shows the results of differential display comparing cDNAs of several patient DCIS samples with cDNA obtained from normal breast epithelial cells and an early invasive cancer. Although few genes shown in this Figure are differentially expressed in the majority of samples with DCIS, the heterogeneity of gene expression in patient samples is seen.

The differential display method (Liang et al, 1992a and 1992b) allows simultaneous comparison of multiple tissue samples. Initial studies using this method (reverse transcriptase followed by PCR) were unsatisfactory because of unwanted amplification of contaminating DNA in tissue samples and the small size of many of the fragments identified by display. To circumvent some of these problems, we have attempted to combine the advantages of cDNA library screening with the advantages of differential display by:

- 1) Constructing cDNA libraries from the tissue mRNA samples;
- Performing differential display on the plasmid DNA prepared from the cDNA libraries;
- 3) Subcloning the fragments identified by differential display;
- 4) Using the subcloned fragment as a probe to clone the cDNA from the appropriate library.

Example 5

Identification of a gene (RibRed) which is differentially expressed in multiple

NCDCIS cases

Employing these methods, 10 differentially expressed clones were identified and the seven that showed the greatest difference in expression between multiple samples were further characterized by DNA sequencing. Comparison of the sequenced clones with GenBank demonstrated that six of the clones are apparently unique sequences (although further DNA sequencing is necessary); but that one of the clones (here termed DCIS-1 and described in Sequence Listing No. 1) showed 90% homology to the

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previously cloned hamster gene encoding the M2 subunit of ribonucleotide reductase (Pavloff et al, 1992; Hurta et al, 1991; Hurta et al, 1991). Although human M2 ribonucleotide reductase has been cloned previously, comparison of the hamster cDNA sequence with our clone and with the prior human clone indicates that DCIS-1 is homologous to an alternatively poly-adenylated form of the human ribonucleotide reductase which has not been cloned previously. Figure 6 shows a comparison of the sequence between DCIS-1 and the human and hamster genes.

Because of our concern that different patients may have differential gene expression which is idiosyncratic (or related to morphological differences in biopsy appearance) and not necessarily related to the induction or progression of DCIS, we simultaneously analyzed gene expression in multiple DCIS samples compared to multiple control samples. We constructed cDNA libraries from the following samples:

- 1) Cultured HMEC epithelial cells;
- 2) Reduction mammoplasty: 11 year old with virginal hyperplasia;
- 3) Reduction mammoplasty: 28 year old patient;
- 4) Reduction mammoplasty: 35 year old patient;
- 5) DCIS patient #12;
- 6) DCIS patient #8;
- 7) DCIS patient #10;
- 8) DCIS patient #10 from an area of invasive cancer adjacent to DCIS;

In addition to the samples we employed to construct cDNA libraries shown above, we also obtained frozen tissue samples from 7 more DCIS patients, 2 cellular fibroadenoma samples, and samples of "usual hyperplasia" and atypical hyperplasia.

Because the DCIS clones were identified by cloning methods which include selection and amplification, it was important to confirm by nuclease protection assays that the genes were differentially expressed in the original unselected, unamplified RNA samples (Figure 7).

This approach allowed identification of a human gene similar to the hamster RibRed gene (coding for the M2 subunit) and 7 other human genes as genes which are differentially expressed in a majority of cases of DCIS in human breast tissue. The table of differentially expressed genes lists the genes which have been identified as

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differentially expressed genes in DCIS tissue samples as compared to that in normal tissue (Figure 9).

EXAMPLE 6

Methods for studying potential use of differentially expressed genes for diagnostic screening

One advantage of the differential display method is that it allows comparison of multiple tissue samples of pre-invasive or invasive breast cancer. For example, use of this method has successfully demonstrated that the M2 subunit ribonucleotide reductase gene is differentially expressed in 4 out of 5 pre-invasive breast cancer tissue samples. It is significant that the M2 subunit is involved in the regulation of the ribonucleotide reductase gene and is found to be over-expressed in abnormal tissue samples.

Identification of differentially expressed genes may lead to discovery of genes which are potentially useful for breast cancer screening. Of particular interest are genes whose expression is restricted to breast epithelial cells and whose gene products are secreted. Screening for secreted proteins is possible by using the known hydrophobic sequences which encode leader sequences as one primer for differential display. The identification of secreted proteins which are specific for early breast premalignancy (or even early invasive cancer) would provide an important tool for early breast cancer screening programs. If a differentially expressed gene has not been cloned previously (or if details of its expression are unknown or uncertain) then nuclease protection assays or Northern blots can be performed on RNA prepared from tissue samples from a variety of tissues to determine if expression of this gene is restricted to breast. If necessary cDNA libraries prepared from other tissues can be added to the differential display screen as a way to identify only those genes which are expressed in early breast cancer and, in addition, are only expressed in breast tissue.

Once differentially expressed genes have been initially characterized for expression in pre-malignant and malignant breast disease, antibodies to the protein products of potentially useful genes can be developed and employed for immunohistochemistry (Harlow et al, 1988). This will provide an additional test to determine whether the expression of this gene is restricted to the breast. Subsequently, these antibodies will

be used to detect the presence of this protein present in the blood of patients with preinvasive and/or invasive cancer. By assaying for serum protein levels in the same patients who exhibited elevated expression of the gene in their tissue samples it will be possible to determine whether a gene product is being secreted into the blood.

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EXAMPLE 7

Decreased expression of BRCA1 accelerates growth and is observed during breast cancer progression

Breast cancer occurs in hereditary and sporadic forms. Recently the BRCA 1 gene has been cloned and shown to be mutated in kindreds with hereditary breast and ovarian cancer (Hall et al. 1990, Miki, Y. et al. 1994, Friedman et al. 1994, Castilla et al. 1994, Simard et al. 1994). Although 92% of families with two or more cases of early-onset breast cancer and two cases of ovarian cancer have germ-line mutations in BRCA 1 (Narod et al. in press), the gene has not been shown to be mutated in any truly sporadic case to date (Futreal et al. 1994). Despite the surprising paucity of somatically acquired mutations in sporadic breast cancer, it is still a likely tumor suppressor gene with a key role in breast epithelial cell biology. The BRCA 1 gene encodes a protein of 1863 amino acids with a predicted zinc finger domain observed in proteins which regulate gene transcription.

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As an initial characterization of the regulation and function of the BRCA 1 gene, we analyzed and manipulated expression of BRCA 1 mRNA levels. The results taken together indicate that the BRCA 1 gene product is a negative regulator of mammary cell proliferation which is expressed at diminished levels in sporadic breast cancer.

Expression of BRCA1 mRNA during breast cancer progression

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As described above, microscopy-directed cloning has been employed to compare gene expression in normal mammary epithelium, carcinoma in-situ, and invasive breast cancer. This method produces predominantly epithelial mRNA with minimal contamination from stromal elements and we used this approach to obtain mRNA from normal neoplastic tissues from patients without a family history of breast cancer. Expression of BRCA1 exon 24 in human breast tissue samples is shown in Fig. 1. The legend of Fig. 1 is as follows.

The following tissue samples were used for mRNA isolation: Normal tissue samples: NL1-cultured human breast epithelial cells, NL2- Histologically normal breast tissue from an 11 year old undergoing a reduction mammoplasty, NL4- histologically normal breast tissue from an 14 year old undergoing a reduction mammoplasty. Carcinoma-in-situ samples are #6, #8, #10, #12, #23 (comedo type), #41, #55; and invasive cancer samples #10CA (invasive cancer from the same patient with carcinoma-in-situ), 36CA, 1CA. All of these tissue samples were obtained from patients who had no family history of hereditary breast cancer and RNA preparation was performed as described above.

PCR detection of BRCA1 exon 24 in cDNA libraries from the following tissue samples is described in Figure 10A. Lane 1: human genomic DNA, lane 2: NL1, lane 3: NL4, lane 4: \$8, lane 5: #12, lane 6: #10, lane 7: #10CA, lane 8: #41, lane 9: #23, lane 10: 36CA, lane 11: lambda DNA. The arrow points to the expected 113 bp band.

Nuclease protection assays of microdissected mRNA from tissue samples are described in Fig. 10B. One ug of mRNA from each tissue sample was hybridized with 32P-labelled, T7 polymerase-generated RNA probes for BRCA1 and human glyceraldehyde-3-phosphate dehydrogenase (GAPD) which produce expected protected fragments of 113 and 140 respectively as indicated by the lines on the right. Data were quantitated by phosphorimaging. The hybridizing intensity of each BRCA1 band was normalized to its respective GAPD band. The normalized values of NL1, NL2, and NL4 were intensity in each sample relative to 1. Sample 1 employs human leukocyte mRNA; Samples 2-4 are NL1, NL2, and NL4; Samples 5-9 are #6(2.8), 8(3.7), 10(2.8), 12 (5.9), and 55 (1.4); and 10-12 are #10CA (0.07), 36CA (0.13), and 1CA (0.2).

Fig. 10 shows that BRCA1 exon 24 mRNA is expressed at 5-10 fold higher levels in normal mammary tissue than in invasive breast cancer samples. Initial studies showed detectable levels of BRCA1 cDNA in a cDNA library prepared from a tissue sample with preinvasive carcinoma-in-situ but not in normal breast cancer invasive breast cancer cDNA libraries (Figure 10A). Because this method is relatively insensitive we directly quantitated BRCA1 mRNA by nuclease protection assays in RNA samples obtained by our microdissection method described above. These assays

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epithelial tissue (lanes 2-4, Figure 10B) is 5-15 fold higher than that in breast cancer (lanes 10-12, Figure 10B). The highest levels of BRCA1 are observed in samples from non-comedo ductal carcinoma-in-situ (lanes 5-9, Figure 10B), a premalignant breast lesion with a finite, but relatively low rate of progression to invasion (Betsill et at., 1978, Page, D.L. et al., 1982, Page and Dupont, 1990).

Because these studies suggested that invasive breast cancer exhibited lower mRNA levels than normal breast epithelial cells, we compared expression of paired samples of normal breast and invasive cancer from the same patient (Figure 11A; compare lanes 2 and 3, 4 and 5, 6 and 7). The legend of Fig. 11 is as follows.

Nuclease protection assays of RNA obtained from paired samples of invasive breast cancer and histologically normal breast tissue are shown in Fig. 11A. Samples in lanes 2 and 3 (first patient), 4 and 5 (second patient), 6 and 7 (third patient) are from invasive cancer and normal breast tissue respectively. Lane 1 is NL1 mRNA as described in legend to Fig. 10 and lane 8 is human leukocyte mRNA. Ratios of BRCA1/GAPD for each sample: lane 1: 25.9, lane 2: 1.8, lane 3: 7.6, lane 4: 2.0, lane 5: 12.4, lane 6: 0.7, lane 7: 6.0. The probes and methods are as described in Fig. 10 except the GAPD probe was of lower specific activity to improve quantitation.

Nuclease protection assays of RNA from a series of invasive breast cancer tissue samples (lanes 2-9 compared with NL1 (lane 1) and leukocyte mRNA (lane 10) are shown in Fig. 11B. Ratios of BRCA1/GAPD for each sample: lane 1: 19.1, lane 2: 0.3, lane 3: 1.8, lane 4: 1.6, lane 5: 0.2, lane 6: 0.3, lane 7: 1.9, lane 8: 0, lane 9: 0.6.

Although the samples were paired in Fig. 11A, they were not microdissected so this approach overestimates the relative expression level of invasive samples because they have a greater percentage of epithelial cells. RNA levels were four to eight fold higher in samples derived from normal breast than in samples derived from invasive breast cancer. We next analyzed expression levels in 8 non-hereditary invasive cancer samples (Figure 11B: lanes 2-7). Although these samples showed some variability in expression level, all had lower levels of BRCA1 mRNA (determined by ratio of

BRCA1 to GAPD) than the primary breast epithelial cell line or the normal breast samples shown in Figure 11A.

Effects of BRCA1 gene inhibition on proliferative rate and gene expression

Having demonstrated that mRNA expression levels of BRCA1 are higher in normal mammary cells than in cancer cells, we used antisense methods to test the hypothesis that BRCA1 expression inhibits cell growth. Unmodified 18 base deoxyribonucleotide complementary to the BRCA1 translation initiation site were synthesized and added to cultures of primary mammary epithelial cells (Stampfer et al. 1980) or MCF-7 breast cancer cells (Soule and McGrath, 1980). Figure 12 is graph showing growth rate of human primary mammary epithelial cells (A), MCF-7 cells (B), retinal pigmented epithelial cells (C), cultured as described below. Points and bars represent the mean and the 95% confidence interval of triplicate counts of cells incubated with a single bolus of the indicated concentration of antisense or control sense deoxyribonucleotide.

The morphologic appearance of the cell lines was not noticeably changed by addition of antisense oligonucleotide, but the proliferative rate was faster. Incubation of cells with 40 uM anti-BRCA1 oligonucleotide produced accelerated growth of both normal (Figure 12A) and malignant mammary cells (Figure 12B), but did not affect the growth of human retinal pigmented epithelial cells (Figure 12C). An intermediate dose of anti-BRCA1 oligonucleotide produced a less pronounced but significant increase in cell growth rate. This was not a toxic effect of the oligonucleotide since a control "sense" oligomer with the same GC content did not increase the proliferation rate, and because an addition of a 10 fold excess of sense oligomer to the anti-BRCA1 oligomer reversed the growth activation.

In order to critically evaluate the function of BRCA1 gene inhibition on growth stimulation and cell cycle progression it was necessary to identify a gene whose expression is cell cycle regulated in human mammary cells. The gene encoding the M2 subunit of ribonucleotide reductase is amplified in conditions of nucleotide starvation (Hurta and Wright 1992) and as shown above, exhibits elevated levels of expression in premalignant breast disease. Because ribonucleotide reductase constitutes the rate limiting step in DNA synthesis, we reasoned that it might be cell cycle regulated in a

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synchronous growth model such as MCF-7 cells which can be growth arrested by tamoxifen and then restimulated by estrogen (Aitken et al. 1985, Arteaga et al. 1989). MCF-7 cells were growth arrested by tamoxifen for 48 hours and then stimulated at time zero (0) with 1uM estradiol (+E) or control vehicle (-E). Inhibition of DNA synthesis by tamoxifen and induction of synthesis by estrogen were confirmed by nuclear labelling studies with tritiated thymidine.

Fig. 13 panels A and B show that transcription of the ribonucleotide reductase M2 gene is cell cycle regulated, inhibited by tamoxifen, and induced by estrogen. Fig. 13A is a Northern blot of mRNA from synchronized MCF-7 cells. At the indicated time in hours, total cellular RNA was isolated and Northern blotting performed using the 1.6 Kb Eco RI fragment from our cloned human ribonucleotide reductase cDNA described above. Two mRNA species of 1.6 and 3.4 Kb are observed in these studies.

Fig. 13B shows nuclear runon studies of synchronized MCF-7 cells were performed by our published methods (Holt et al 1988) employing the 1.6 Kb fragment of ribonucleotide reductase described above (RR); the 1.8 Kb fragment of Topoisomerase II (Topo) described in the Olsen et al. 1993); the 1.0 Kb cyclophilingene (Thompson et al. 1994) used as a constitutive control; and 18S ribosomal RNA (Thompson et al. 1994). Con represents cells which were grown for 48 hours but not treated with tamoxifen.

Antisense inhibition is a useful strategy for studying gene expression which is dependent on expression of the antisense target gene (Robinson-Benion and Holt, in press, 1995), e.g. genes whose expression is directly or indirectly dependent on BRCA1 levels. Fig. 14 demonstrates that antisense inhibition of BRCA1 results in a corresponding increased expression of M2 ribonucleotide reductase mRNA. A nuclease protection assay of mRNA derived from primary mammary epithelial cells (lanes 1-4, 9-10) or MCF-7 cells (lanes 5-8, 11-12) cultured for 4 days with antisense or control oligonucleotide was performed under the following conditions: no oligonucleotide (lanes 1 and 5); 40uM antiBRCA1 (lanes 2,6,10,12); 4uM antiBRCA1 (lanes 3 and 7); 40uM sense control (lanes 4,8,9,11). Probes for BRCA1 and GAPD are as described for

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Figure 10, and the ribonucleotide reductase M2 probe (RR) detects the 200 bp probe is described above.

Ribonucleotide reductase mRNA levels are highest in samples treated with 40 uM anti-BRCA1 oligonucleotide for both primary mammary epithelial cells and for MCF-7 cells (Fig. 14). Antisense inhibition of BRCA1 results in a 70-90% inhibition of mRNA levels in anti-BRCA1 treated cells compared with cells treated with the "sense" control oligonucleotide (compare lanes 9 and 10, Fig. 14). Note that MCF-7 cells have lower levels of BRCA1 than the normal mammary epithelial cells (compare lanes 1 and 5, Fig. 14) anti-BRCA 1 since the antisense inhibition may drop BRCA1 levels below a critical threshold which normally functions to inhibit growth.

Methodology

Tissue samples. Freshly obtained breast biopsy or reduction mammoplasty specimens were frozen and then RNA was obtained following the microdissection method described above. Lesions were selected which were microlocalized and homogenous so that pure lesions could be obtained by 2 mm punches. Samples which had admixed normal epithelial, carcinoma-in-situ, or invasive cancer were not used for this study. Family history was obtained by chart review and/or interview to exclude familial breast cancer cases.

Nuclease Protection Assays. PCR primers were derived from BRCA1 U14680); forward GenBank (Accession number sequence in CAATTGGGCAGATGTGT 3' and reverse 5' CTGGGGGATCTGGGGTATCA 3' which amplify a 113 bp region from exon 24, corresponding to bases 5587 to 5699 of the human BRCA1. This region was selected because this exon has not been reported to be differentially spliced unlike more 5' exons. The BRCA1 probe was cloned by subcloning this 113 bp band from normal human genomic DNA into PCRscriptSK and screening for correct orientation. One ug of mRNA from each tissue sample was hybridized with 32P-labelled, T7 polymerase-generated RNA probes for BRCA1 and human glyceraldehyde-3-phosphate dehydrogenase (GADP) which would produce expected protected fragments of 113 and 140 respectively. The construction and use of the GADP probe for RNA standardization has been described above. The probe for

ribonucleotide reductase M2 mRNA is the same as above and detects a 200 bp protected fragment.

Antisense oligonucleotide studies. Unmodified deoxyribonucleotide were analyzed by gel electrophoresis and UV shadowing and shown to be homogenous and of appropriate size. These oligonucleotide were purified by multiple lyophilization and solubilized in buffered media as described (Holt et al. 1988). Sequence of the unmodified antiBRCA1 oligonucleotide 5' AAGAGCAGATAAATCCAT 3' and the complementary sense oligonucleotide 5' ATGGATTTATCTGCTCTT 3' correspond to the presumed translation initiation site at bases 12-137 of the GenBank sequence. The antisense oligonucleotide sequence was searched against Genbank and no significant homologies were identified to genes except BRCA1. Oligonucleotides were used according to our published methods (Holt et al. 1988). Primary mammary epithelial cells were cultured in serum-free medium supplemented with epidermal growth factor, insulin, hydrocortisone, ethanolamine, phosphorylethanolamine, and bovine pituitary extract. MCF-7 cells were cultured in Minimum Essential Medium Eagle (Modified) with Earle's salts and 2g/L sodium bicarbonate m supplemented with 2mM Lglutamine, GMS-A (Gibco Cat. #680-1300AD), nonessential amino acids, and 2.5% fetal calf serum. Retinal pigmented perithelial cells were cultured in DMEM and 10% calf serum.

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Our results indicate that the BRCA1 gene is expressed at higher levels in normal mammary cells than in breast cancer cells and that diminished expression of BRCA1 increased the proliferative rate of breast cells. This correlates well with the recent finding that patients with BRCA1 gene-linked hereditary breast cancer have tumors that grow more rapidly than comparable sporadic tumors (Marcus, J. et al. 1994). The decreased mRNA levels which were observed in sporadic breast cancers are not a consequence of differential splicing of the gene since the RNAs were quantitated with probes from the 3' end of the mRNA which is not a region where differential splicing is reported to occur (Miki, Y. et al 1994). Invasive sporadic cancers have BRCA1 mRNA levels which vary from 0 (in one case) to 20% of the levels observed in normal human mammary epithelium.

Examples 8 and 9 describe applications of the discovery of the function of the BRCA1 gene. Example 8 describes a gene therapy method and example 9 describes a drug screening method. The discovery of the diminished expression of the BRCA1 mRNA in breast cancer using the microdissection techniques of this invention provides an important scientific basis for these examples.

Example 8

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Gene Therapy method based on determination of the function of the BRCA1 Gene

Viral vectors containing a DNA sequence that codes for a protein having an amino acid sequence as essentially set forth in SEQ ID NO:49 can be constructed using techniques that are well known in the art. This sequence includes the BRCA1 gene product. Viral vectors containing a DNA sequence essentially as set forth in SEQ ID NO:47 (the BRCA1 gene) can be also constructed using techniques that are well known in the art. Retroviral vectors, adenoviral vectors, or adeno-associated viral vectors are all useful methods for delivering genes into breast cancer cells. An excellent candidate for use in breast cancer gene therapy is a Moloney-based retroviral vector with a breast selective MMTV promoter which we have reported previously (Wong et al). The viral vector is constructed by cloning the DNA sequence essentially as set forth in SEQ ID:47 into a retroviral vector such as a breast selective vector. Most preferably, the full-length (coding region) cDNA for BRCA1 is cloned into the retroviral vector. The retroviral vector would then be transfected into virus producing cells in the following manner: Viruses are prepared by transfecting PA317 cells with retroviral vector DNAs which were purified as described in Wong et al. Following transfection, the PA317 cells are split and then treated with G418 until individual clones can be identified and expanded. Each clone is then screened for its titer by analyzing its ability to transfer G418 resistance (since the retroviral vector contains a Neomycin resistance gene). The clones which have the highest titer are then frozen in numerous aliquots and tested for sterility, presence of replication-competent retrovirus, and presence of mycoplasm. The methods generally employed for construction and production of retroviral vectors have been described in Muller, 1990.

Once high titer viral vector producing clones have been identified, then patients with breast cancer can be treated by the following protocol: Viral vector expressing

BRCA1 is infused into either solid tumors or infused into malignant effusions as a means for altering the growth of the tumor (since it is shown above that the BRCA1 gene product decreases the growth rate of breast cancer cells). Because viral vectors can efficiently transduce a high percentage of cancer cells, the tumors would be growth inhibited.

Example 9

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Method of Screening Compounds Capable of Activating Promoter Region of the BRCA1 Gene

The discovery of the function of the BRCA1 gene provides a clear utility in that induction of expression of the gene and the resulting increase in level of protein encoded by the gene in the breast cancer cell should slow the proliferation of the breast cancer cells. Induction of expression of the gene can be caused by administering a compound to a patient that stimulates the regulatory regions of this gene, such as the promoter.

A method for screening compounds that activate the promoter of the BRCA1 gene is designed in the following way. A promoter sequence is a DNA segment that upregulates the expression of a gene. A sequence essentially as set forth in SEQ ID NO:48 can be ligated into a suitable vector, such as a plasmid, that contains a reporter gene using standard recombinant DNA techniques of restriction enzyme digests, ligation of fragment into vector, and transformation of bacteria. SEQ ID NO:48 includes the

promoter sequence of the BRCA1 gene. A reporter gene is a gene that produces a readily detectable product. Examples of appropriate reporter genes which could be employed for this purpose include Beta-galactosidase or the chloramphenicol

acetyltransferase gene.

The BRCA1 promoter/reporter gene combination can then be cloned into an expression vector or viral vector by standard recombinant DNA methods. Breast cancer cells can then be transfected with the expression vector containing the BRCA1 promoter/reporter gene using standard transfection methods which we have reported previously (Holt et al. PNAS 1986). A stable transformant with appropriate low level expression (breast cancer cells have low level BRCA1 expression as shown above) will be identified and then characterized to demonstrate proper DNA integration and

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expression. Methods of establishing and characterizing stable transformants have been described (Holt. MCB, 1994). Once an appropriate stable transformant cell line is identified, then we can plate the cell line in a manner than permits screening of hundreds or thousands of drugs or biological agents (for example in multiple 96 well microtiter plates). Level of expression of the reporter gene can be quantitated and agents which activate expression are thus identified. A positive result (i.e. induction of the promoter region) results in increased levels of the reporter gene resulting in either an increase in color (Beta-galactosidase assay) or specific radioactivity (Chloramphenicol acetyltransferase activity) through a reaction between the protein encoded by the reporter gene and a compound in the reaction medium. The compound produced by the reaction between the reporter gene protein and the compound in the reaction medium is the cause of the increase in color or specific radioactivity. These compounds can be called indicator compounds in that their presence indicates that the drug or biologial agent activitated the BRCA1 promoter. Methods for standardizing and performing Beta-galactosidase or chloramphenicol acetyltransferase assays have been reported (Holt et. al. MCB 1994). This method would be useful for initial screening of agents which increase BRCA1 expression. These agents could then be tested in more rigorous assays of breast cancer growth such as nude mouse tumor assays (Arteaga et al). This approach allows mass screening of large numbers of agents, sparing more rigorous animal tests for only promising compounds which score in the reporter gene assay described herein.

Thus, although there have been described particular embodiments of the present invention of a new and useful "Method for Detection and Treatment of Breast Cancer", it is not intended that such embodiments be construed as limitations upon the scope of this invention except as set forth in the following claims. It will be apparent to those skilled in the art that many changes and modifications may be made without departing from the invention in its broader aspects. For example, the above described techniques may be used in the diagnosis of other diseases and detection of differential genetic expression from microscopically-directed tissue samples of pathologic tissue. The production of a cDNA library produced as a result of the differential expression of genes in pathologic tissue in comparison to normal tissue provides the opportunity for

further adiagnostic capabilities. Further, although there have been described certain experimental conditions used in the preferred embodiment, it is not intended that such conditions be construed as limitations upon the scope of this invention except as set forth in the claims.

The following references are included to provide details of scientific technology herein incorporated by reference to the extent that they provide additional information for the purposes of indicating the background of the invention or illustrating the state of the art.

REFERENCES

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ADDITIONAL DESCRIPTION OF THE FIGURES

Figure 2: Model for premalignant conditions, highlighting magnitude of risk for progression to clinical malignancy. Terms from human breast neoplasia are used: no proliferative disease (No Pro), proliferative disease without alypia (PDWA), typical hyperplasia (AH), carcinoma in situ (CIS). As is proposal of tumor progression each stage is more likely to proceed to the next (dotted lines), but could also remain stable (horizontal lines, probably fairly frequent), or directly proceed to develop a clone of cells with malignant behavior (vertical lines, becoming more likely further to right.)

Figure 5: Differential display of cDNAs obtained from patient tissue samples and controls. Rescued cDNA library samples were used as templates for low stringency PCR with the primers 5'GATGAGTTCGTGTCCGTACAACTGG3' and 5' GGTTATCGAAATCAGCCACAGCGCC3'; 40 cycles were performed at conditions described above. Samples (See legend to Figure 4): Lane 1 - #12; Lanes 2 and 3: separate phage rescues of NL1 to show reproducibility of the assay; Lane 4 - #8; Lane 5 - #10; Lane 6 - #10CA; Lane 7 - control from the rescued phage vector without cDNA inserts. Arrows mark cDNAs which are overexpressed in DCIS versus normal. Arrowheads mark cDNAs which are differentially expressed in the invasive cancer (note this may reflect contamination from stromal cells). The bar marks a cDNA which is expressed in normal breast cells at higher levels than in DCIS or invasive cancer.

Figure 7: Expression of DCIS-1 mRNA in tissue mRNA samples analyzed by RNase protection assay. Probes: GADH probe and DCIS-1 clone probe which was generated by linearizing the rescued plasmid with Pvu II and should generate a 200 bp protected fragment. RNA samples were labeled as in the legend to Figure 4.

)

SEQUENCE LISTINGS

(1)	GENERAL	INFORMATION:	•
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- (i) APPLICANT: HOLT, JEFFREY T.
 - JENSEN, ROY A.

PAGE, DAVID L.

OBERMILLER, PATRICE S.

ROBINSON-BENION, CHERYL L.

THOMPSON, MARILYN E.

- (ii) TITLE OF INVENTION: METHOD FOR DETECTION AND TREATMENTS OF BREAST CANCER
- (iii) NUMBER OF SEQUENCES: 49
- (iv) CORRESPONDENCE ADDRESS:
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 - (B) STREET: 27TH FLOOR, L & C TOWER, 401 CHURCH
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- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Diskette, 3.50 inch, 800 kB storage
 - (B) COMPUTER: IBM PC/XT/AT compatible
 - (C) OPERATING SYSTEM: MS-DOS (version 5.0)
 - (D) SOFTWARE: WordPerfect 5.1/WordPerfect Editor
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: U.S. 08/182,961
 - (B) FILING DATE: 14 JAN 1994

(viii)	ATTO	RNEY/AGENT INFORMATION:
	(A)	NAME: I.C. WADDEY, JR.
	(B)	REGISTRATION NUMBER: 25,180
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(ix)	TELE	COMMUNICATION INFORMATION (O):
	(A)	TELEPHONE: (615) 242-2400
	(B)	TELEFAX: (615) 242-2221
	(C)	TELEX:
	(2)	INFORMATION FOR SEQ ID NO:1:
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	(C)	STRANDEDNESS: double
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(iii)	HYP	OTHETICAL: no
(iv)	ANT	I-SENSE: no
(v)	ORIO	GINAL SOURCE
	(A)	ORGANISM: Homo sapiens sapiens
	(C)	INDIVIDUAL/ISOLATE: sample of non-comedo DCIS
	(D)	DEVELOPMENTAL STAGE: adult
	(F)	TISSUE TYPE: female breast
	(G)	CELL TYPE: ductal carcinoma in situ
	(H)	CELL LINE: not derived from a cell line
	(I)	ORGANELLE: no
(vii)	IMIN	MEDIATE SOURCE:
	(A)	LIBRARY: cDNA library derived from human
	(B)	CLONE: obtained from identification of differential
		gene expression

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(D)	OTHER INFORMATION: gene encoding M2 subunit of			
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PUBI	LICATION INFORMATION: unpublished			
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ANT	ANTI-SENSE: no			
ORI	GINAL SOURCE			
ORG	ANISM: Homo sapiens sapiens			
(C)	INDIVIDUAL/ISOLATE: sample of non-comedo DCIS			
(D)	DEVELOPMENTAL STAGE: adult			
(F)	TISSUE TYPE: female breast			
	(A) (B) (C) FEAT (A) (B) (C) (D) PUBI (K) SEQUE (A) (B) (C) (D) MOI HYP ANT ORIC ORG (C) (D)			

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	(H)	CELL LINE: not derived from a cell line		
	(I)	ORGANELLE: no		
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	(B)	CLONE: obtained from identification of differential gene		
expression				
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	(C)	IDENTIFICATION METHOD: microscopically-directed		
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(x)	PUBL	ICATION INFORMATION: unpublished		
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(iii)	HYPOTHETICAL: no			
(iv)	ANTI-SENSE: no			
(v)	ORIG	INAL SOURCE		
	(A)	ORGANISM: Homo sapiens sapiens		

	(C)	INDIVIDUAL/ISOLATE: Sample of holl-contedo Dello
	(D)	DEVELOPMENTAL STAGE: adult
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	(H)	CELL LINE: not derived from a cell line
	(I)	ORGANELLE: no
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expression		
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	(C)	UNITS: unknown
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	(A)	NAME/KEY: DCIS-3
	(B)	
	(C)	IDENTIFICATION METHOD: microscopically-directed
	-	oling and differential display
(x)	PUB	LICATION INFORMATION: unpublished
	(K)	
(xi)	SEQU	JENCE DESCRIPTION: SEQ ID NO: 3:
TGCCCGATGT GT	_	ACTIGGCGCTG TGGCTGATTT CGATAA 46 INFORMATION FOR SEQ ID NO:4:
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(iii)		
(iv)	AN	TI-SENSE: no

(v)	ORIG	INAL SOURCE		
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	(C)	INDIVIDUAL/ISOLATE: sample of non-comedo DCIS		
	(D)	DEVELOPMENTAL STAGE: adult		
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	(H)	CELL LINE: not derived from a cell line		
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expression				
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(iv)	ANTI-SENSE: no		
(v)	ORIG	INAL SOURCE	
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•	(C)	INDIVIDUAL/ISOLATE: sample of non-comedo DCIS	
	(D)	DEVELOPMENTAL STAGE: adult	
	(F)	TISSUE TYPE: female breast	
	(G)	CELL TYPE: ductal carcinoma in situ	
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	(I)	ORGANELLE: no	
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	(B)	CLONE: obtained from identification of differential gene	
expression			
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	(B)	MAP POSITION: unknown	
	(C)	UNITS: unknown	
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(x)	PUBI	LICATION INFORMATION: unpublished	
	(K)	RELEVANT RESIDUES IN SEQ ID NO: 5	
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(x)	PUBLICATION INFORMATION: unpublished		
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(iii)	HYPO	OTHETICAL: no	
(iv)	ANTI	-SENSE: no	
(v)	ORIG	INAL SOURCE	
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	(C)	INDIVIDUAL/ISOLATE: sample of non-comedo DCIS	
	(D)	DEVELOPMENTAL STAGE: adult	
	(F)	TISSUE TYPE: female breast	
	(G)	CELL TYPE: ductal carcinoma in situ	
	(H)	CELL LINE: not derived from a cell line	
	(T)	ORGANELLE: no	
(vii)	IMM	EDIATE SOURCE:	
	(A)	LIBRARY: cDNA library derived from human	
	(B)	CLONE: obtained rom identification of differential gene	
expression			
(viii)	POSI	TION IN GENOME:	
	(A)	CHROMOSOME/SEGMENT: unknown	
	(B)	MAP POSITION: unknown	
	(C)	UNITS: unknown	
(ix)	FEA'	TURE:	
	(A)	NAME/KEY: DCIS-7	
	(B)	LOCATION: L27643	

	(C)	IDENTIFICATION	METHOD:	microscopically-directed
	sampling and differential display			
(x)	PUBLICATION INFORMATION: unpublished			ned
	(K)	RELEVANT RESIDU	ES IN SEQ ID	NO: 7
(xi)	SEQU	ENCE DESCRIPTION	: SEQ ID NO:	7:
		CCCGCGCC CCCCCCTCCG TCGGAAT		
ATCCATAGGA TGTGC	GAGTTA GT (2)	itigii INFORMATION FOR	8:SEO ID NO	-
(i)	` ,	ENCE CHARACTERIS	_	
(*)	_	LENGTH: 25		
	• ,	TYPE: nucleic acid		
	, ,	STRANDEDNESS: si	ngle	
		TOPOLOGY: linear	C	
(ii)	, ,	CULE TYPE: DNA		
	(A)	DESCRIPTION: PCR	primer	
(iii)	НҮРО	THETICAL: yes		
(iv)	ANTI-	SENSE: no		
(v)	FRAG	MENT TYPE: oligonu	cleotide	
(xi)	SEQU	ENCE DESCRIPTION	: SEQ ID NO:	8:
CGCGACGGCC GCGC	GTCTGC CA	GGG 25		
	(2)	INFORMATION FOR	R SEQ ID NO:	9
(i)	SEQU	ENCE CHARACTERI	STICS:	
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: si	ingle	
	(D) ·	TOPOLOGY: linear		
(ii)	MOLI	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR	t primer	
(iii)	HYPO	OTHETICAL: yes		
(iv)	ANTI	-SENSE: no		

(v)	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQUE	ENCE DESCRIPTION: SEQ ID NO: 9:	
CGCCCCTGCG TTA	CCCTCCC C	GCCG 25	
	(2)	INFORMATION FOR SEQ ID NO:10	
(i)	SEQU	ENCE CHARACTERISTICS:	
	(A)	LENGTH: 25	
	(B)	TYPE: nucleic acid	
	(C)	STRANDEDNESS: single	
	(D)	TOPOLOGY: linear	
(ii)	MOL	ECULE TYPE: DNA	
	(A)	DESCRIPTION: PCR primer	
(iii)	HYPO	OTHETICAL: yes	
(iv)	ANT	I-SENSE: no	
(v)	FRAC	GMENT TYPE: oligonucleotide	
(xi)	SEQU	JENCE DESCRIPTION: SEQ ID NO: 10:	
GGATGGCGTC CTC	STAACCCG A	CGCT 25	
	(2)	INFORMATION FOR SEQ ID NO:11	
(i)	SEQ	JENCE CHARACTERISTICS:	
	(A)	LENGTH: 25	
	(B)	TYPE: nucleic acid	
	(C)	STRANDEDNESS: single	
	(D)	TOPOLOGY: linear	
(ii)	MOL	ECULE TYPE: DNA	
	(A)	DESCRIPTION: PCR primer	
(iii)	HYP	OTHETICAL: yes	
(iv)	ANTI-SENSE: no		
(v)	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 11:	
ACTGGGCTGT CC	TGCGGTGG	ceece 25	
	(2)	THEODMATION FOR SEC ID NO.12	

(i)	SEQU	ENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOL	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYPO	OTHETICAL: yes		
(iv)	ANTI	-SENSE: no		
(v)	FRAC	GMENT TYPE: oligonucleotide		
(xi)	SEQU	JENCE DESCRIPTION: SEQ ID NO: 12:		
CTGAGAGGTA G	CCGCGCGGA G	GCTG 25		
	(2)	INFORMATION FOR SEQ ID NO:13		
(i)	SEQU	JENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
		STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOL	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYP	OTHETICAL: yes		
(iv)	ANTI-SENSE: no			
(v)	FRA	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQU	SEQUENCE DESCRIPTION: SEQ ID NO: 13:		
GCCTGGCCGC G	ACACGGATT A	accgc 25		
	(2)	INFORMATION FOR SEQ ID NO:14		
(i)	SEQ	JENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
•	(C)	STRANDEDNESS: single		

	` ,	TOPOLOGY: linear		
(ii)	MOLECULE TYPE: DNA			
	(A)	DESCRIPTION: PCR primer		
(iii)	HYPOTHETICAL: yes			
(iv)	ANTI	ANTI-SENSE: no		
(v)	FRAGMENT TYPE: oligonucleotide			
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 14:			
TTAGCGCATG GTG	GACCTGG A	GACG 25		
	(2)	INFORMATION FOR SEQ ID NO:15		
(i)	SEQU	JENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
•	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOL	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYPOTHETICAL: yes			
(iv)	ANTI-SENSE: no			
(v)	FRAGMENT TYPE: oligonucleotide			
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 15			
TGTGGTTACG TC	AGCGAAGG '	TAATA 25		
	(2)	INFORMATION FOR SEQ ID NO:16		
(i)	SEQ	UENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOI	LECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYPOTHETICAL: yes			
(iv)	ANTI-SENSE: no			

(v)	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 16:		
AGTEGEAEGE ATGT	CACGCT CO	egcc 25	
	(2)	INFORMATION FOR SEQ ID NO:17	
(i)	SEQU	ENCE CHARACTERISTICS:	
·	(A)	LENGTH: 25	
	(B)	TYPE: nucleic acid	
	(C)	STRANDEDNESS: single	
	(D)	TOPOLOGY: linear	
(ii)	MOLI	ECULE TYPE: DNA	
	(A)	DESCRIPTION: PCR primer	
(iii)	HYPO	OTHETICAL: yes	
(iv)	ANTI-SENSE: no		
(v)	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 17		
TATCCAAGCG GCAG	GCTACG A	GGCC 25	
	(2)	INFORMATION FOR SEQ ID NO:18	
(i)	SEQU	JENCE CHARACTERISTICS:	
	(A)	LENGTH: 25	
	(B)	TYPE: nucleic acid	
	(C)	STRANDEDNESS: single	
	(D)	TOPOLOGY: linear	
(ii)	MOL	ECULE TYPE: DNA	
	(A)	DESCRIPTION: PCR primer	
(iii)	HYP	OTHETICAL: yes	
(iv)	ANTI-SENSE: no		
(v)	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 18		
GGCGCGCCCG ACG	STCTGGT A	ITCTA 25	
	(2)	INFORMATION FOR SEQ ID NO:19	
	-		

(i)	SEQU	SEQUENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOL	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYP	OTHETICAL: yes		
(iv)	ANT	ANTI-SENSE: no		
(v)	FRA	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQ	SEQUENCE DESCRIPTION: SEQ ID NO: 19		
CTCCCTCCCC GC	SACTEGGGG 1	TAGT 25		
	(2)	INFORMATION FOR SEQ ID NO:20		
(i)	SEQ	UENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOL	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYP	HYPOTHETICAL: yes		
(iv)	ANT	ANTI-SENSE: no		
(v)	FRA	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQ	SEQUENCE DESCRIPTION: SEQ ID NO: 20		
ATGCGGGCGG C	TCGGGCCTG	GTCGC 25		
	(2)	INFORMATION FOR SEQ ID NO:21		
(i)	SEQ	UENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
ď	(C)	STRANDEDNESS: single		
	(D)	TOPOI OGV: linear		

(ii)	MOL	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYPO	OTHETICAL: yes		
(iv)	ANTI	ANTI-SENSE: no		
(v)	FRAC	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQU	TENCE DESCRIPTION: SEQ ID NO: 21:		
CGTGAAGCCT A	TGCCCTCCC T	CAAC 25		
	(2)	INFORMATION FOR SEQ ID NO:22		
(i)	SEQU	JENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOL	ECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYP	OTHETICAL: yes		
(iv)	ANT	ANTI-SENSE: no		
(v)	FRA	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQU	ENCE DESCRIPTION: SEQ ID NO: 22:		
GTGCCGTCGT	AGCCCTTCAG (CGATC 25		
	(2)	INFORMATION FOR SEQ ID NO:23		
(i)	SEQ	UENCE CHARACTERISTICS:		
	(A)	LENGTH: 25		
	(B)	TYPE: nucleic acid		
	(C)	STRANDEDNESS: single		
	(D)	TOPOLOGY: linear		
(ii)	MOI	LECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer		
(iii)	HYF	HYPOTHETICAL: yes		
(iv)	AN	ANTI-SENSE: no		
(v)	FRAGMENT TYPE: oligonucleotide			

(xi)	SEQU	TENCE DESCRIPTION: SEQ ID NO: 23:	
GCGACACTAG GCTC	CCGGAG G	AGGG 25	
	(2)	INFORMATION FOR SEQ ID NO:24	
(i)	SEQUENCE CHARACTERISTICS:		
	(A)	LENGTH: 25	
•	(B)	TYPE: nucleic acid	
	(C)	STRANDEDNESS: single	
•	(D)	TOPOLOGY: linear	
(ii)	MOLECULE TYPE: DNA		
	(A)	DESCRIPTION: PCR primer	
(iii)	HYPO	OTHETICAL: yes	
(iv)	ANTI-SENSE: no		
(v)	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 24:		
TGGGCCAGGC CTCC	eeeccc e	GTAT 25	
	(2)	INFORMATION FOR SEQ ID NO:25	
(i)	SEQU	ENCE CHARACTERISTICS:	
	(A)	LENGTH: 25	
	(B)	TYPE: nucleic acid	
	(C)	STRANDEDNESS: single	
	(D)	TOPOLOGY: linear	
(ii)	MOL	ECULE TYPE: DNA	
·	(A)	DESCRIPTION: PCR primer	
(iii)	HYPO	OTHETICAL: yes	
(iv)	ANTI-SENSE: no		
(v)	FRAGMENT TYPE: oligonucleotide		
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 25:		
CCGGAACTGC GATAGCGTCC GTCCC 25			
	(2)	INFORMATION FOR SEQ ID NO:26	

(i)	SEQU	SEQUENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	STRANDEDNESS: single						
	(D)	TOPOLOGY: linear						
(ii)	MOL	ECULE TYPE: DNA						
	(A)	DESCRIPTION: PCR primer						
(iii)	HYP	OTHETICAL: yes						
(iv)	ANT	-SENSE: no						
(v)	FRAC	GMENT TYPE: oligonucleotide						
(xi)	SEQU	JENCE DESCRIPTION: SEQ ID NO: 26:						
AGCGGACACC T	GTTTCCCGA G	AGCC 25						
	(2)	INFORMATION FOR SEQ ID NO:27						
(i)	SEQ	JENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	STRANDEDNESS: single						
	(D)	TOPOLOGY: linear						
(ii)	MOL	ECULE TYPE: DNA						
	(A)	DESCRIPTION: PCR primer						
(iii)	HYP	OTHETICAL: yes						
(iv)	ANT	I-SENSE: no						
(v)	FRA	GMENT TYPE: oligonucleotide						
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 27:						
AACGGGTGGA	CATCCGCCTG	ccgcc 25						
	(2)	INFORMATION FOR SEQ ID NO:28						
(i)	SEQ	UENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	STRANDEDNESS: single						
	W)	TOPOLOGY: linear						

(ii)	MOLE	ECULE TYPE: DNA								
	(A)	DESCRIPTION: PCR primer								
(iii)	HYPO	OTHETICAL: yes								
(iv)	ANTI-SENSE: no									
(v)	FRAC	MENT TYPE: oligonucleotide								
(xi)	SEQU	ENCE DESCRIPTION: SEQ ID NO: 28:								
TGAACCACGA TGTC	AATCGT C	CCGA 25								
	(2)	INFORMATION FOR SEQ ID NO:29								
(i)	SEQU	JENCE CHARACTERISTICS:								
•	(A)	LENGTH: 25								
	(B)	TYPE: nucleic acid								
	(C)	STRANDEDNESS: single								
	(D)	TOPOLOGY: linear								
(ii)	MOL	ECULE TYPE: DNA								
	(A)	DESCRIPTION: PCR primer								
(iii)	HYP	OTHETICAL: yes								
(iv)	ANT	I-SENSE: no								
(v)	FRA	GMENT TYPE: oligonucleotide								
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 29								
TCATCCCCGC CG/	MAGACGC	TEGEC 25								
	(2)	INFORMATION FOR SEQ ID NO:30								
(i)	SEQ	UENCE CHARACTERISTICS:								
	(A)	LENGTH: 25								
	(B)	TYPE: nucleic acid								
	(C)	STRANDEDNESS: single								
	(D)	TOPOLOGY: linear								
(ii)	MO	LECULE TYPE: DNA								
	(A)	DESCRIPTION: PCR primer								
(iii)	HY	POTHETICAL: yes								
(iv)	AN	Π-SENSE: no								
(v)	FRA	AGMENT TYPE: oligonucleotide								

(xi)	SEQU	TENCE DESCRIPTION: SEQ ID NO: 30:
ATAGGETGEG GEAC	GCGCTG G	GACT 25
	(2)	INFORMATION FOR SEQ ID NO:31
(i)	SEQU	JENCE CHARACTERISTICS:
	(A)	LENGTH: 25
	(B)	TYPE: nucleic acid
	(C)	STRANDEDNESS: single
	(D)	TOPOLOGY: linear
(ii)	MOL	ECULE TYPE: DNA
	(A)	DESCRIPTION: PCR primer
(iii)	HYPO	OTHETICAL: yes
(iv)	ANT	I-SENSE: no
(v)	FRAC	GMENT TYPE: oligonucleotide
(xi)	SEQU	JENCE DESCRIPTION: SEQ ID NO: 31:
GACCAGGTGC GCA	CGAGCAT G	TACA 25
	(2)	INFORMATION FOR SEQ ID NO:32
(i)	SEQU	JENCE CHARACTERISTICS:
	(A)	LENGTH: 25
	(B)	TYPE: nucleic acid
	(C)	STRANDEDNESS: single
	(D)	TOPOLOGY: linear
(ii)	MOL	ECULE TYPE: DNA
	(A)	DESCRIPTION: PCR primer
(iii)	HYP	OTHETICAL: yes
(iv)	ANT	I-SENSE: no
(v)	FRA	GMENT TYPE: oligonucleotide
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 32:
AGCGTAGTCA TCG	GCCTTCG	
	(2)	INFORMATION FOR SEQ ID NO:33

(i)	SEQU	SEQUENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	STRANDEDNESS: single						
	(D)	TOPOLOGY: linear						
(ii)	MOL	ECULE TYPE: DNA						
	(A)	DESCRIPTION: PCR primer						
(iii)	HYP	OTHETICAL: yes						
(iv)	ANT	I-SENSE: no						
(v)	FRA	GMENT TYPE: oligonucleotide						
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 33:						
GGCCCCTAGC	CCAGGGTGAA (GCCCA 25						
	(2)	INFORMATION FOR SEQ ID NO:34						
(i)	SEQ	SEQUENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	STRANDEDNESS: single						
	(D)	TOPOLOGY: linear						
(ii)	MOI	LECULE TYPE: DNA						
	(A)	DESCRIPTION: PCR primer						
(iii)	HYF	POTHETICAL: yes						
(iv)	ANT	T-SENSE: no						
(v)	FRA	GMENT TYPE: oligonucleotide						
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 34						
CCCAGTGCTA	CGGGCCGCCC	CAAGC 25						
	(2)	INFORMATION FOR SEQ ID NO:35						
(i)	SEQ	UENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	CTD ANDEDNESS: single						

	(D) TOPOLOGY: linear									
(ii)	MOLECULE TYPE: DNA									
	(A) DESCRIPTION: PCR primer									
(iii)	HYPOTHETICAL: yes									
(iv)	ANTI-SENSE: no									
(v)	FRAGMENT TYPE: oligonucleotide									
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 35									
CCTTCCTGGG T	TACCTGCCC TCGGG 25									
·	(2) INFORMATION FOR SEQ ID NO:36									
(i)	SEQUENCE CHARACTERISTICS:									
	(A) LENGTH: 25									
	(B) TYPE: nucleic acid									
	(C) STRANDEDNESS: single									
	(D) TOPOLOGY: linear									
(ii)	MOLECULE TYPE: DNA									
	(A) DESCRIPTION: PCR primer									
(iii)	HYPOTHETICAL: yes									
(iv)	ANTI-SENSE: no									
(v)	FRAGMENT TYPE: oligonucleotide									
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 30									
TCCGGACAGC	GCCACGCCA AGGGC 25									
	(2) INFORMATION FOR SEQ ID NO:37									
(i)	SEQUENCE CHARACTERISTICS:									
	(A) LENGTH: 25									
	(B) TYPE: nucleic acid									
	(C) STRANDEDNESS: single									
	(D) TOPOLOGY: linear									
(ii)	MOLECULE TYPE: DNA									
	(A) DESCRIPTION: PCR primer									
(iii)	HYPOTHETICAL: yes									
(iv)	ANTI-SENSE: no									

(v)	FRAGMENT TYPE: oligonucleotide							
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 37:							
ACGCGCTGGT CCACC	GAGGC CT	GAT 25						
	(2)	INFORMATION FOR SEQ ID NO:38						
(i)	SEQU	ENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	STRANDEDNESS: single						
	(D)	TOPOLOGY: linear						
(ii)	MOLE	ECULE TYPE: DNA						
	(A)	DESCRIPTION: PCR primer						
(iii)	HYPC	OTHETICAL: yes						
(iv)	ANTI-SENSE: no							
(v)	FRAGMENT TYPE: oligonucleotide							
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 38:							
CGATGCAAGG CCAC	CAGCAC T	CGAC 25						
	(2)	INFORMATION FOR SEQ ID NO:39						
(i)	SEQU	JENCE CHARACTERISTICS:						
	(A)	LENGTH: 25						
	(B)	TYPE: nucleic acid						
	(C)	STRANDEDNESS: single						
	(D)	TOPOLOGY: linear						
(ii)	MOL	ECULE TYPE: DNA						
	(A)	DESCRIPTION: PCR primer						
(iii)	HYP	OTHETICAL: yes						
(iv)	ANT	I-SENSE: no						
(v)	FRA	GMENT TYPE: oligonucleotide						
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 39						
CCCCCGGAGC GG/	ACCACCGG	ACGTG 25						

(2) INFORMATION FOR SEQ ID NO:40

(i)	SEQU	ENCE CHARACTERISTICS:					
	(A)	LENGTH: 25					
	(B)	TYPE: nucleic acid					
	(C)	STRANDEDNESS: single					
	(D)	TOPOLOGY: linear					
(ii)	MOL	ECULE TYPE: DNA					
	(A)	DESCRIPTION: PCR primer					
(iii)	HYP	OTHETICAL: yes					
(iv)	ANT	I-SENSE: no					
(v)	FRA	GMENT TYPE: oligonucleotide					
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 40:					
AGCGGGGAGG	GATCGGGGGC	CAAGC 25					
	(2)	INFORMATION FOR SEQ ID NO:41					
(i)	• •	UENCE CHARACTERISTICS:					
(-)		LENGTH: 25					
		TYPE: nucleic acid					
		STRANDEDNESS: single					
		TOPOLOGY: linear					
(ii)	MO	LECULE TYPE: DNA					
	(A)	DESCRIPTION: PCR primer					
(iii)	HY	POTHETICAL: yes					
(iv)	AN	II-SENSE: no					
(v)	FRA	AGMENT TYPE: oligonucleotide					
(xi)	SEC	QUENCE DESCRIPTION: SEQ ID NO: 41:					
GCCTGGTG	TA GGCAGGCAGC	TCTTA 25					
	(2)	INFORMATION FOR SEQ ID NO:42					
(i)	SEC	QUENCE CHARACTERISTICS:					
	(A)	LENGTH: 25					
	(B)	TYPE: nucleic acid					
	(C)	STRANDEDNESS: single					
	(D)	TOPOLOGY: linear					

(ii)	MOLE	CULE TYPE: DNA								
	(A)	DESCRIPTION: PCR primer								
(iii)	HYPC	THETICAL: yes								
(iv)	ANTI-	ANTI-SENSE: no								
(v)	FRAG	MENT TYPE: oligonucleotide								
(xi)	SEQU	ENCE DESCRIPTION: SEQ ID NO: 42:								
CCACCCCTGT A	STGCGGGCT G	CGAG 25								
	(2)	INFORMATION FOR SEQ ID NO:43								
(i)	SEQU	JENCE CHARACTERISTICS:								
	(A)	LENGTH: 25								
	(B)	TYPE: nucleic acid								
	(C)	STRANDEDNESS: single								
	(D)	TOPOLOGY: linear								
(ii)	MOL	ECULE TYPE: DNA								
	(A)	DESCRIPTION: PCR primer								
(iii)	HYP	OTHETICAL: yes								
(iv)	ANT	I-SENSE: no								
(v)	FRA	GMENT TYPE: oligonucleotide								
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 43:								
GGAACCCGAC	GCCCGTCCAG	GGTTC 25								
	(2)	INFORMATION FOR SEQ ID NO:44								
(i)	SEQ	UENCE CHARACTERISTICS:								
	(A)	LENGTH: 25								
	(B)	TYPE: nucleic acid								
	(C)	STRANDEDNESS: single								
	(D)	TOPOLOGY: linear								
(ii)	MO	LECULE TYPE: DNA								
	(A)	DESCRIPTION: PCR primer								
(iii)	HYI	POTHETICAL: yes								
(iv)	AN	Π-SENSE: no								

(v)	FRAC	SMENT TYPE: oligonucleotide
(xi)	SEQU	TENCE DESCRIPTION: SEQ ID NO: 44:
TCGGGCAGCA AGGC	CGGGAC G	CTCC 25
	(2)	INFORMATION FOR SEQ ID NO:45
(i)	SEQU	JENCE CHARACTERISTICS:
	(A)	LENGTH: 25
	(B)	TYPE: nucleic acid
	(C)	STRANDEDNESS: single
	(D)	TOPOLOGY: linear
(ii)	MOL	ECULE TYPE: DNA
	(A)	DESCRIPTION: PCR primer
(iii)	HYP	OTHETICAL: yes
(iv)	ANT	I-SENSE: no
(v)	FRA	GMENT TYPE: oligonucleotide
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 45:
GACGGGGGAC GGG	CTAGGTG (GCTTA 25
	(2)	INFORMATION FOR SEQ ID NO:46
(i)	SEQ	UENCE CHARACTERISTICS:
	(A)	LENGTH: 25
	(B)	TYPE: nucleic acid
	(C)	STRANDEDNESS: single
	(D)	TOPOLOGY: linear
(ii)	MOI	LECULE TYPE: DNA
	(A)	DESCRIPTION: PCR primer
(iii)	HYF	POTHETICAL: yes
(iv)	AN	TI-SENSE: no
(v)	FRA	GMENT TYPE: oligonucleotide
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO: 46
CTTGTTGCCG GC	GGAGAGGG	CIGCC 25
	(2)	INFORMATION FOR SEQ ID NO:47:

(i)	SEQU	JENCE CHARACTERISTICS:							
	(A)	LENGTH: 5712							
	(B)	TYPE: nucleic acid							
	(C)	STRANDEDNESS: double							
	(D)	TOPOLOGY: linear							
(ii)	MOL	ECULE TYPE: cDNA to mRNA							
(iii)	HYPO	OTHETICAL: no							
(iv)	ANT	-SENSE: no							
(v)	ORIC	INAL SOURCE							
	(A)	ORGANISM: Homo sapiens sapiens							
	(C)	INDIVIDUAL/ISOLATE:							
	(D)	DEVELOPMENTAL STAGE: adult							
	(F)	TISSUE TYPE: female breast							
	(G)	CELL TYPE: ductal carcinoma in situ, invasive breast cancer							
		and normal breast tissue							
	(H)	CELL LINE: not derived from a cell line							
•	(I)	ORGANELLE: no							
(vii)	IMM	EDIATE SOURCE:							
	(A)	LIBRARY: cDNA library derived from human							
	(B)	CLONE: obtained using published sequence							
(viii)	POSI	TION IN GENOME:							
	(A)	CHROMOSOME/SEGMENT: unknown							
	(B)	MAP POSITION: unknown							
	(C)	UNITS: unknown							
(ix)	FEA'	TURE:							
	(A)	NAME/KEY: BRCA1							
	(B)	LOCATION: GenBank accession no. U14680							
	(C)	IDENTIFICATION METHOD: microscopically-directed							
		sampling and nuclease protection assay							
	(D)	OTHER INFORMATION: gene encoding BRCA1 protein							

(x)	PUBL	ICATION INFOR	RMATION:	
	(A)	AUTHORS: Mi	iki, Y., et. al.	
	(B)	TITLE: A	strong candidate ge	ne for the breast and ovarian
			ncer susceptibility	
	(0)		-	50.10 21(0111)
	(C)	JOURNAL: Sci		
	(D)	VOLUME: 266	i	
	(E)	PAGES: 66-71		
	(F)	DATE: 1994		
	(K)	RELEVANT RE	ESIDUES IN SEQ I	D NO: 47
(xi)	SEQU	ENCE DESCRIP	TION: SEQ ID N	O:47:
agctcgctga ga	acttcctgg 8	cccgcacc aggctgtggg	gtttctcaga taactgggcc	60
			ttcattggaa cagaaagaa	119
		cgc gtt gaa gaa gta		167
Met Asp Leu :	ser Ala Leu 5	Arg Val Glu Glu Val	15	
•		gag tgt ccc atc tgt		215
		Glu Cys Pro Ile Cys		
	20	25	30	
		tgt gac cac ata ttt		263
	Ser Thr Lys	Cys Asp His Ile Phe	Cys Lys Phe Cys Met	
Stg ass ctt	CTC BBC CBO	40 aag aaa ggg cct tca		311
		Lys Lys Gly Pro Ser		
50		55	60	
		agg agc cta caa gas		359
		Arg Ser Leu Gin Glu	Ser Thr Arg Phe Ser 80	
65	70	75 ttg aaa atc att tgt		407
			Ala Phe Gin Leu Asp	
	85	90	95	
aca ggt ttg	gag tat gca	aac agc tat aat ttt	gca aaa aag gaa aat	455
Thr Gly Leu	Glu Tyr Ala		Ala Lys Lys Glu Asn	
•	100	105	110	503
			atc atc caa agt atg	303
115	GIO RIS LEG	120	125	
	aac cgt gco		agt gaa ccc gaa aat	551
			ser Glu Pro Glu Asn	
130		135	140	
			e ctc tct aac ctt gga	5 99
			n Leu Ser Asn Leu Gly 5 160	
145	15	D 15!	, 160	

cact	gtg	aga	act	ctg	egg	808	aag	Cag	cgg	ata	Caa	cct	CBB	888	acg	647
Thr																
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tct	at c	***	•••		***		***	nat	***	tet	088	det	acc	att	aat	695
Ser	-															0,2
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-	210					215					220					
gct	tgt	988	ttt	tct	989	acg	gat	atg	aca	aat	act	988	cat	cat	CBB	839
Ala	Cys	Glu	Phe	Ser	Glu	Thr	Asp	Val	Thr	Asn	Thr	Glu	His	His	Gin	
225					230					235					240	
ccc	agt	786	aat	gat	ttg	880	acc	act	gag	889	cgt	gca	gct	gag	agg	887
Pro	Ser	Asn	Asn	Asp	Leu	Asn	Thr	Thr	Glu	Lys	Arg	Ala	Ale	Glu	Arg	
				245					250					255		
cat	CCB	988	aag	tat	gao	ggt	agt	tct	gtt	tca	880	ttg	cat	gtg	989	935
His	Pro	Glu	Lys	Tyr	Gln	Gly	Ser	Ser	Val	Ser	Asn	Leu	His	Val	Glu	
			260					265					270			
CCB	tgt	ggc	808	aat	act	cat	gcc	agc	tca	tta	cag	cat	989	вас	agc	983
Pro	Cys	Gly	Thr	Asn	Thr	His	Ala	Ser	Ser	Leu	Gln	His	Glu	Asn	Ser	
		275					280				•	285				
agt	tta	tta	ctc	act	888	gac	ega	atg	aat	gta	988	889	gct	gae	ttc	1031
Ser	Leu	Leu	Leu	Thr	Lys	Asp	Arg	Het	Asn	Val	Glu	Lys	Ala	Glu	Phe	
	290					295					300					
tgt	aat	886	agc	888	свд	cct	ggc	tta	gca	899	agc	CBB	cat	8 B C	898	1079
Cys	Asn	Lys	Ser	Lys	Gln	Pro	Gly	Leu	Alo	Arg	Ser	Gln	His	Asn	Arg	
305					310					315					320	
tgg	gct	gga	agt	889	gaa	aca	tgt	aat	gat	a 99	cgg	act	ccc	agc	aca	1127
Тгр	Ala	Gly	Ser	Lys	Glu	Thr	Cys	Asn	Asp	Arg	Arg	Thr	Pro	Ser	Thr	
				325					330					335		
gaa	888	889	gta	gat	ctg	aat	gct	gat	ccc	ctg	tgt	989	898	886	gaa	1175
															Glu	
			340					345					350			
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															Glu	
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gat	att			ata	aca	cta	aat	BQC	890	att	Cag	888	gtt	881	gag	1271
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	370					375					380					
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				420					425					430			
q	cc	egt	gat	cct	cat	gag	gct	tta	ata	tgt	888	agt	gaa	aga	gtt	CBC	1463
					His												
-		-	435					440		-			445				
				ata	gag	aat	aat	att	caa	gac	888	ata	ttt	999	888	BCC	1511
				-	Glu	-											
		450	JC.	•••	•••		455		•••		-,-	460		,			
					gca	200		ccc	886	tta	age		ata	act	BAD	aat	1559
					Ala												
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				515					520					525			
•	cct	gas	atg	ata	aat	cag	998	act	880	CBB	acg	gag	CB9	aat	ggt	CBB	1751
•	Pro	Glu	Met	Ile	Asn	Gln	Gly	Thr	Asn	Gln	Thr	Glu	Gln	Asn	Gly	Gln	
			530					535					540				
•	gtg	atg	aat	att	act	aat	agt	ggt	cat	989	aat	888	aca	888	ggt	gat	1799
,	Val	Het	Asn	Ile	Thr	Asn	Ser	Gly	His	Glu	Asn	Lys	Thr	Lys	Gly	Asp	
		545					550					555					
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	560					565					570					575	
	gaa	tct	gct	ttc	888	acg	888	gct	gaa	cct	ata	agc	agc	agt	ata	agc	1895
	Glu	Ser	Ale	Phe	Lys	Thr	Lys	Ala	Glu	Pro	ile	Ser	Ser	Ser	116	Ser	
					580					585					590		
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gca	cc	t	ggt	tct	ttt	act	889	tgt	tca	aat	BCC	agt	860	ctt	888	gaa	2279
Ala	Pr	0	Gly	Ser	Phe	Thr	Lys	Cys	Ser	Asn	The	Ser	Glu	Leu	Lys	Glu	
	70						710					715					
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												Glu					
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	G 1	t	888	gtg	tct	aat	aat	gct	вае	gac	ccc	888	gat	ctc	atg	tta	2375
												Lys					
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												Val					
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		865					87		_ •-						a ca	a agt	2807
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				Glu													
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ace	at1	ago	cgt:	aat	880	att	898	986	881	t gt1	ttt	886	988	e gc	C B	gc	3239
Thr	- 11	Ser	Arg	Asr	Asr	110	Arg	GL	J Ast	n Vai	l Pho	ty:	s Gl	u Al	a S	er	
	102	25				103	0				103	35					
				t aat													3287
Se	r Se	r Ası	n Ile	e Asr	n Glu	اaV د	Gly	/ Se	r Se	r Th	r Asi	n Gli	u Va	l Gl			
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Se	r Il	e As	n Gl	u 11	e Gl	y Se	r Se	r As	p Gl	u As	n Ii	e Gl	n Al			.eu	
				10						65					70		3383
99	t ag	88 8	c ag	a 99	g cc	88 6	e tt	g 88	t gc	t at	g ct	t ag	B TT	a gg	99 5	/ L	2263
Gl	y Ar	g As		g Gl	у Рг	o Ly	s Le			a Me	t Le	NU AF		085	, y		
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tt	g ca	- CC	t ge	u Va	C T8		. CI		,, ,,	41 Pr	o Gi	v Se	;. A:	sn C	vs I	Lys	
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			1	155				1	160				1	1165			
8	8C 8	tt e	8 9	8 86	gt t	ct g	ct g	tt t	tt e	gc E	888	gc g	tc (cag	888	gga	367
A	sp 1	le L	ys 0	ilu S	er S	er A	le V	al F	he s	Ser t	Lys :			Gln	Lys	Gly	
		1	170				1	175				•	1180				

989	ctt	agc	899	agt	cct	agc	cct	ttc	BCC	cat	aca	cat	ttg	gct	cag	3719
					Pro											
	1185		_			1190					1195					
ggt	tac	cga	898	999	gcc	aag	888	tta	gag	tcc	tca	988	gag	aac	tta	3767
					Ala											
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		QBG	gat	gaa	gag	ctt	ccc	tgc	ttc	CBB	cac	ttg	tta	ttt	ggt	3815
					Glu											
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888	gta	880	aat	ete	cct	tct	cag	tct	act	agg	cat	agc	acc	gtt	gct	3863
Lys	Val	Asn	Asn	lle	Pro	Ser	Gln	Ser	Thr	Arg	His	Ser	Thr	Val	Ala	
			123	5				1240)				124	5		
acc	gag	tgt	ctg	tct	888	BBC	808	gag	988	aat	tta	tta	tca	ttg	889	3911
					Lys											
		125	0				125	5				126	0			
aat	890	tta	ast	gac	tgc	agt	asc	cag	gta	ata	ttg	gca	889	gca	tct	3959
					Cys											
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CAG	989	cat	cac	ctt	agt	989	gaa	aca	888	tgt	tct	gct	agc	ttg	ttt	4007
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Asp Asp	Arg	Тгр	Туг	Met	His	Ser	Cys	Ser	Gly	Ser	Leu	Gln	Asn	Arg	
1505				1510)				1515	5				1520	
aac tac	CC8	tct	CBB	gag	989	ctc	att	aag	gtt	gtt	gat	gtg	gag	989	4727
Asn Tyr	Pro	Pro	Gln	Glu	Glu	Leu	Ile	Lys	Vel	Val	Asp	Val	Glu	Glu	
			1525	5				1530)				1535		
caa cag	ctg	988	989	tct	888	CCB	CBC	gat	ttg	ecg	988	808	tct	tac	4775
Gln Gln				Ser	Gly	Pro			Leu	Thr	Glu			Туг	
		1540					1545					1550			
ttg cca			_							_	_		-		4823
Leu Pro	_		Asp	Leu	Glu			Pro	Туг	Leu		_	Gly	Ile	
	1555					1560	ס				156	5			
agc ctc			_	-		-		-			-	_	_	-	4871
Ser Leu		Ser	Asp	Asp			Ser	Asp	Pro			Asp	Arg	Ala	
157	-				157					1580	-				
cca gag			-	-				•							4919
Pro Glu	5er	Ala	Arg		-	Asn	He	Pro			Thr	Ser	ALB		
1585				159					159					1600	1047
asa gtt			-												4967
Lys Val	PLO	GIN			Val	AIB	Glu			GIN	2er	Pro			
			160					161					1615		ENTE
gct cat			_												5015
Ala His	INF		•	INF	ALB	ыу	•		ALB	met	GIU			Val	
		1620					162					163			5047
agc agg															5063
Ser Arg			Pro	Glu	Fen			Ser	inr	Glu			ASN	Lys	
	1635					164					164	-			F 4 4 4
aga atg		_													5111
Arg Met		met	vat	VAL		_	r en	INC	PFO			rne	net	ren	
165					165					166					E150
gtg tac	_		-												5159
Val Tyr	LYS	Phe	ALB			HIS	HIS	11e			INC	ASN	FEU	1680	
1665				167					167						E 202
act gas															5207
Thr Glu	Glu	זחר	Thr		Val	VBl		Lys		ASP	ALB	610	1 PNE		

															***	5255
									gga Gly							3233
Lys	GIU	Ary	1700		Lys	',	7 1.0	1709		•••		٠.,	1710			
ata	att	agc			tgg	gtg	acc	cag	tct	att	888	gaa	898	886	atg	5303
-	-								Ser							
		1715					172					172	_			
ctg	aat	989	cat	gat	ttt	988	gtc	aga	998	gat	gtg	gtc	aat	998	aga	5351
Leu	Asn	Glu	His	Asp	Phe	Glu	Val	Arg	Gly	Asp	Val	Val	Asn	Gly	Arg	
	173					173					174	-				
									988							5399
		Gln	Gly	Pro			Ala	Arg	Glu	5er 175		ASP	Arg	Lys	1760	
174					175		***	***	999			800	825	ata		5447
									Gly							•
FILE	AI 9	Gty	260	176		•,5	-,-	.,.	177					177	_	
aca	gat	CBB	ctg			atg	gta	cag	ctg	tgt	ggt	gct	tct	gtg	gtg	5495
									Leu							
			178	0				1785	,				179	0		
_									808							5543
Lys	Glu	Leu	Ser	Ser	Phe	Thr	Leu	Gly	Thr	Gly	Val			Ile	Val	
		179	-				180			_		180	-			EE01
									gac							5591
Val	Va (Pro	ASP	ALE	181			3 ASP	ASI	182				ile	
900			tat	GBC	908		-	gte	3 800	: cge			gts	ttg	gac	5639
															Asp	
182			,		183					183					1840	
															CCC	5687
Ser	Val	Ale	Leu	ı Tyr	Glr	Cys	Gli	n Gli	u Lec	J Asp) The	Ty	r Leu	_	e Pro	
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Glr	110	e Pro			. 111	s iyı	r									-
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(i))			S	EQ'	UE	NC	E C	HA	RA	CT.	ERI	STI	CS	:	
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				(C)	S	TR	AN	DE	DN	ES!	S: d	oub	le		
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(ii	i)			N	101	LEC	UI	E	ГҮР	E:	DN	Αı	regu	lato	ory se	quence
	ii)			H	IYF	TO	HE	TIC	CAL	.: ne	o					
(i				A	נאו	T-S	EN	SE:	no							

(v)	ORIG	INAL SOURCE
	(A)	ORGANISM: Homo sapiens sapiens
	(C)	INDIVIDUAL/ISOLATE:
	(D)	DEVELOPMENTAL STAGE: adult
	(F)	TISSUE TYPE: female breast
	(G)	CELL TYPE: normal breast
	(H)	CELL LINE: not derived from a cell line
	(I)	ORGANELLE: no
(vii)	IMM	EDIATE SOURCE:
	(A)	LIBRARY: cDNA library derived from human
	(B)	CLONE: obtained using published sequence
(viii)	POSI	TION IN GENOME:
	(A)	CHROMOSOME/SEGMENT: unknown
	(B)	MAP POSITION: unknown
	(C)	UNITS: unknown
(ix)	FEAT	TURE:
	(A)	NAME/KEY: BRCA1 promoter
	(B)	LOCATION:
	(C)	IDENTIFICATION METHOD: restriction enzyme digest
	(D)	OTHER INFORMATION: DNA sequence regulating gene
	enco	ding BRCA1 protein
(x)	PUB	LICATION INFORMATION:
	(A)	AUTHORS: Brown et al.
	(B)	TITLE: Scientific Correspondence
	(C)	JOURNAL: Nature
	(D)	VOLUME: 372
	(E)	PAGES: 733
	(F)	DATE: 22/29 DECEMBER 1994
	(K)	RELEVANT RESIDUES IN SEQ ID NO: 48
(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO:48:
TTCCGGGAC	T CTACTACCTT	TACCCAGACG AGAGGGTGAA GGCCTCCTGA TCGCAGGGGC 60

CCAGTTATCT GAGAAACCCC ACAGCCTGGT GCGGGGTCCA GGAAGTCTCA GCGAGCTCAC 120

CELL LINE: not derived from a cell line

ORGANELLE: no

NAME/KEY: BRCA1 protein

GCCGCGCAGT	CGCAGTTTTA	ATTTATCTGT AATTCCCGCG CTTTTCCGTT GCCACGGAAA 180
CCAAGGGGCT	ACCGCTAAGC	AGCAGCCTCT CAGAATACGA AATCAAGGTA CAATCAGAGG 240
AAGGGAGGGA	CAGAAAGAGC	CAAGCGTCTC TCGGGGCTCT GGATTGGCCA CCCAGTCTGC 300
CCCCGGATGA	CGTAAAAGGA	AAGAGACGGA AGAGGAAGAA TTCTACCTGA GTTCGCCGTA 360
		CTACGCTTCC AGTTGCGGCT TATTACGTCA CAGTAATTGC 420
		CACCTGAGGC CTGAATATCA GCGTAAGATA GTGTCCAAAG 480
		ATTACCCCAC TCTTTCCGCC CTAATGGAGT CCTCCAGTTT 540
		GGGAGGTGGA GGGAAAGAAC TACTATTTCC AACATGCATT 600
		CACACTGTTC CTTGGAAACT GTAGTCTTAT GGAGAGGAAC 660
		CAATTCTCAC GGAAATCCAG TGGATAGATT GGAGACCTCC 720
· -		AGTAATATIG GGTTGTTATG TTCTCCTATC TTGAGAGCAG 780
		ATAGGAAGAC TACGATTCCC ATCCAGCCCC ACGAGTCTCG 840
-		CAGTEGCCTG CGGGGACGCA GTGGGCGCCG AATTTGCCTG 900
		TEGTICACATE TECEGRACITEC TAGTITECECE CCTCAGCATE 960 GGGTTCAGGT TECTTCTECE ECECCCATE GACGCAATCT 1020
		GGGTTCAGGT TGCTTCTGCC CCGCCCCATC GACGCAATCT 1020 CGTTTTGAGG GACAAGTGGT GAGAGCCAAT CATCTTGGCG 1080
		ACTAGTTACT GTCTTTATCC GCCATGTTAG ATTCACCCCA 1140
		GTAGCGGACG GTCCTTGCAT TGGCCTCCGG CAGGCGCCCC 1200
		AGGAAGCAGC TGCGGTT 1237
	(2)	INFORMATION FOR SEQ ID NO:49:
	` '	
(i)	SEC	UENCE CHARACTERISTICS:
	(A)	LENGTH: 1863
	(B)	TYPE: amino acid
	(C)	STRANDEDNESS: unknown
	(D)	TOPOLOGY: unknown
(ii)	МО	LECULE TYPE: protein
(iii)	HY	POTHETICAL: no
(iv)	AN	П-SENSE: no
(v)	OR	GINAL SOURCE
	(A)	ORGANISM: Homo sapiens sapiens
	(C)	INDIVIDUAL/ISOLATE:
	(D)	DEVELOPMENTAL STAGE: adult
	(F)	TISSUE TYPE: female breast
	(C)	CELL TYPE: normal breast tissue
	(G)	CELL LIFE. HOUHAI DIEASI USSUE

(H)

(I)

(A)

FEATURE:

(ix)

2			(B)	1	LOC	CAT	OF	1 : :	l to	18	63											
			((C)]	DΕ	NT	IFIC	CAT	ПО	N	ME	TH	OD): o	bse	rva	tion	of	r	nR!	NΑ	and
					á	anti	sens	e in	hib	itio	n o	f BI	RCA	11	gene								
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			1	regu	lato	ory	effe	ct o	n g	rov	⁄th (of h	um	an 1	mam	ıma	ıгу	cell:	s.				
(x)]	PUI	3LI	CA'	ПО	N II	NF	ORI	MA	TIC	N:										
` /			1	(A)		ΑU	TH	ORS	: :	Mik	i, I	۲., ۱	et. a	al.									
				(B)		TIT	LE	:		A s	troi	ng c	and	ida	te ge	ne	for	the	brea	ast	an	d ov	ariaı
										car	cer	sus	сер	tibi	lity	gen	ne F	BRC	A1.				
				(C)		JOI	JRN	IAL	.: S	Scie	nce												
				(D)		vo	LU	ME:	: 2	66													
				(E)		PΑ	GE:	s: 6	56-	71													
				(F)		DA	TE	: 19	994														
				(K)		RE	LE	VAN	17	RE	SID	UE	s II	N S	EQ	ID	NC): 4	9				
(xi)				SE	QUI	EN	CE	DES	SCF	UP	ΠO	N:	SE	Q I	D N	Ю:	49:						
Met /	Asp !	Leu	Ser	Ala	Leu /	Arg '	Val (/al (Gln /	lsn \			Asn								
1 Ala I	Met i	Gin	l Vs	5 Ile	Leu (Glu	Cys !		O Le (Cys !	Leu (Glu I		15 I L e	Lys								
			20				:	25				:	30										
Glu		Val 35	Ser	Thr	Lys		Asp 40	His I	lle	Phe		Lys 45	Phe	Lys	HET								
		Leu	Leu	Asn			Lys	Gly F	Pro			Cys	Pro	Leu	Cys								
	50 Asn	ASD	1 l e	Thr		55 Arg	Ser	Leu (Gln		60 Ser	Thr	Arg	Phe	Ser								
65					70					7 5					80								
Gln	Leu	Val	Glu	Glu 85	Leu	Leu	Lys	lle	1 l e 90	Cys	Ala	Phe	Gin	Leu 95	ASP								
Thr	Gly	Leu	Glu		Ala	Asn	Ser	Tyr .		Phe	Ala	Lys	Lys	Glu	Asn								
		_	100				4	105 Glu	Val	Ser	110	ı le	110 Gln	Ser	Het								
Asn	Ser	Pro 115		HIS	Leu	Lys	120	410	***	Je.	•••	125	•••	•••									
Gly	Туг			Arg	Ala	Lys	Arg	Leu	Leu	Gln	Ser	Glu	Pro	Glu	Asn								
	130					135				- 1-	140	5.00	A c D	Lau	י פוע								
		Leu	Gln	Glu	150		Leu	Ser	val	155		ser	A911	LEU	160								
145 Thr		Ara	Thr	Leu			Lys	Gln	Arg			Pro	Gln	Lys	Thr								
				165					170					175	5								
£	Vo1	Tue	. 114	. Gli	Leu	GI.	Ser	Asp	Ser	Ser	Glu	Asp	Thr	Val	Asn								

190

185

Lys Ale Thr	Tyr Cys	Ser \	al Gl	y Asp	Gln	Glu	Leu		Gln	lle	Thr
195			20					205			
Pro Gin Gly	Thr Arg	Asp (Sto It	e Ser	Leu	Asp	Ser	Ala	Lys	Lys	Ala
210		2	215				220				
Ala Cys Glu	Phe Ser		Thr As	p Val	Thr		Thr	Glu	His		
225		230				235					240
Pro Ser Asn			Asn Th	r Thr		Lys	Arg	Ala			Arg
	245				250					255	
His Pro Glu		Gln	Gly Se		Val	Ser	Asn	Leu		Val	Glu
	260			265	_				270	•	
Pro Cys Gly	Thr Asn	Thr			Ser	Leu	GIN	H15	Glu	ASN	Ser
275	1 a 7h a		28		4.00	Val	ci		Alo	ci	Dho
Ser Leu Leu	Leu in		мър иг 295	g met	ASII	VB	300	Lys	A10	410	riie
290 Cys Asn Lys	San 1.m				A 1 o	4		e la	ui.	Acn	Aca
305	Ser Lys	310	710 60	y Leo	~ 1.0	315	JEI	•	пір	A 611	320
Trp Ala Gly	Serive		Thr Cu	e Aen	Asn		Aro	The	Pro	Ser	
110 210 017	325		,	3 7311	330	~• 5	~- 5	••••		335	••••
Glu Lys Lys			Asn Al	a Asp		Leu	Cvs	Glu	Arg		Glu
,,-	340			345			-,-		350	-•	
Trp Asn Lys	Gin Lys	Leu	Pro Cy	s Ser	Glu	Asn	Pro	Arg	Asp	Thr	Glu
355			36	0				365			
Asp Val Pro	Trp Ile	Thr	Leu As	n Ser	Ser	lle	Gln	Lys	Val	Asn	Glu
370			375				380				
Trp Phe Ser	Arg Ser	Asp	Glu Le	u Leu	Gly	Ser	Asp	Asp	Ser	His	Asp
385		390				395					400
Gly Glu Ser	Glu Ses	Asn	Alo Ly	's Val	Ala	Asp	Val	Leu	Asp	Val	Leu
	40	j			410					415	
Asn Glu Val	Asp Glu	J Tyr	Ser Gl	-		Glu	Lys	He		Leu	Leu
	420			425					430		
Ala Ser Asp	Pro His	Glu			Cys	Lys	Ser			Val	aiH
435			44		_			445			
Ser Lys Ser	Val Gli			e Glu	Asp	Lys			Gly	Lys	Inc
450			455		•	.	460		75-	C 1	
Tyr Arg Lys	LYS AL		Leu Pi	O AST	Leu	5er 475	MIS	VBI	inr	Glu	480
465 Leu Ile Ile	61 A1	470	V-1 C				*1-	11-	CID	clu	
ten ite ite	48		AB1 24	er utt	490			116	um	495	
Pro Leu Thr			Lvc A				Dra	The	Ser		
Pro Lea IIII	50		- yo ~	.g .y.	505			••••	JC .	510	
His Pro Glu			lve l	ve Als			Ale	Val	Gln		
115 PTO GEO	515		_,5 -,	520					525		
Pro Glu Met		n Gin	GLY TI			Thr	Glu	Gln			Gln
530		,		AU. 35				540		- ,	
Val Met Asn		r Asn			: Glu	J AST	LYS	-		Gly	Asp
545		- /= - •	550				555		•	·	-
Ser Ile Gin	Asn Gl	u Lys	-	ro Asi	n Pro	. Ile	Glu	Ser	Leu	Glu	Lys
		,									•

Glu	Ser	alA	Phe	Lys	Thr	Lys	Ala	Glu	Pro	He	Ser	Ser	Ser	Ile	Ser
				580					585					590	
Asn	Glu	Leu	Glu	Leu	Asn	Ιle	Met	His	Asn	Ser	Lys	Ala	Pro	Lys	Lys
			595					600					605		
Asn	Arg	Leu	Arg	Arg	Lys	Ser	Ser	Thr	Arg	His	Ile	His	Als	Leu	Glu
	_	610	•	-	•		615		_			620			
i eu	Val	Val	Ser	Ara	Asn	Ł eu	Ser	Pro	Рго	Asn	Cvs	Thr	Glu	Leu	Gln
	625		•••			630				,,,,,,	635	••••			••••
		5ac	Cve	Sec	Ser		61	61		1 46		l ve	1 200	Tue	Aen
640	vsh	361	Cys	361	645	361	310	3.0	116	650	Lys	Lys	Lys	. , ,	655
		5	·- •			.		•					61	61	
GIN	met	Pro	VBI	_	His	Ser	Arg	ASN		Gin	rea	met	Glu		Lys
	_			660				_	665		_			670	
Glu	Рго	Ala		Gly	Ala	Lys	Lys		Asn	Lys	Pro	Asn		Gln	Thr
			675					680					685		
Ser	Lys	Arg	His	Asp	Ser	Asp	Thr	Phe	Рго	Glu	Leu	Lys	Leu	Thr	Asn
		690					695					700			
Ala	Pro	Gly	Ser	Phe	Thr	Lys	Cys	Ser	Asn	Thr	Ser	Glu	Leu	Lys	Glu
	705					710					715				
Phe	Val	Asn	Pro	Ser	Leu	Pro	Arg	Glu	Glu	Lys	Glu	Glu	Lys	Leu	Glu
720					725					730					735
Thr	Val	Lys	JaV	Ser	Asn	Asn	Ala	Glu	Asp	Pro	Lys	Asp	Leu	Met	Leu
				740					745					750	
Ser	Gly	Glu	Arg	Val	Leu	Gln	Thr	Glu	Arg	Ser	Val	Glu	Ser	Ser	Ser
			755					760					765		
lle	Ser	Leu	Val	Pro	Gly	Thr	Asp	Tyr	GLY	Thr	Gln	Glu	Ser	Ile	Ser
		770			•		775	•	·			780			
i eu	Leu		Vel	Ser	Thr	Leu		I VS	Ala	l ve	Thr		Pro	A sn	l vs
	785			•••		790	01,	-,-		-,-	795				-,-
CVE		Sar	GI n	Cve	Ala		Dha	GI.	Acn	Pro		elv.		11.	Wie
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	Cvc	500	l ve	Acn	Asn	400	Aco	Aco	The		61v	Dhe	l ve	Tvr	
Gty	Cys	361	LYB	820	MBII	~! 9	A511	ASP	825	310	Gty	rne	Lys	830	
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GLU	Ser		Leu	Asp	ALB	GIN		Leu	Gin	ASN	INT		Lys	Val	Ser
	_	850	_				855		•			860			
Lys		Gln	Ser	Phe	Ala			Ser	ASN	Pro		Asn	ALB	Ģlu	GLU
	865					870					875				_
Glu	Cys	Ala	Thr	Phe		Ala	His	Ser	Gly			Lys	Lys	Gln	Ser
880					885					890					895
Рго	Lys	Val	Thr	Phe	Glu	Cys	Glu	Gln	Lys	Glu	Glu	Asn	Gin	Gly	Lys
				900					905					910	
Asn	Glu	Ser	Asn	Ile	Lys	Pro	Val	Gln	Thr	Val	Asn	1 l e	Thr	Ala	Gly
			915					920					925		
Phe	Pro	Val	Val	Gly	Gin	Lys	Asp	Lys	Pro	Val	Asp	Asn	Ala	Lys	Cys
		930					935					940			
Ser	Ile	Lys	Gly	Gly	Ser	Arg	Phe	Cys	Leu	Ser	Ser	Gln	Phe	Arg	Gly
		-	,	,							055				

				uia Clv	LAULEU GIR	Acn
Asn Glu Th	r Gly Leu	Ile Thr P	TO AST LYS		Leu Leu Gin	
960		965		970		9 7 5
Pro Tyr Ar	g Ile Pro	Pro Leu P	he Pro Ile	Lys Ser	Phe Val Lys	Thr
	980		985	i	990	
Lys Cys Ly	s Lys Asn	Leu Leu (ilu Glu Asr	n Phe Glu	Glu His Ser	Met `
-,,,	995		1000		1005	
Ser Pro Cl		Met Gly	lan Glu Asi	n Ile Pro	Ser Thr Val	Ser
	10		1015		1020	
				n Val Phe	Lys Glu Ala	Ser
	I NIS MOII	1030		103		
1025			rly Ser Se		Glu Val Gly	Ser
	in ile Asn		Pri sei se	1050	0.0 (0. 0.,	1055
1040		1045			CI- Ale Chi	
Ser Ile As	sn Glu Ile	Gly Ser			Gin Ale Glu	
	106			65	107	
Gly Arg A	sn Arg Gly	Pro Lys		a Met Leu	Arg Leu Gly	Val
	1075		1080		1085	
Leu Gln P	ro Glu Va	Tyr Lys	Gin Ser Le	Hu Pro Gly	Ser Asn Cys	Lys
1	090		1095		1100	
His Pro G	lu Ile Ly:	s Lys Gln	Glu Tyr Gl	u Glu Val	. Val Gin Thr	Val
1105	•	1110		111		
	sp Phe Se			er Asp Asr	n Leu Glu Glr	1 Pro
1120	٠,٠.٠ و	1125		1130		1135
	er ter Hi		Gin Val C	vs Ser Gli	u Thr Pro Asp	ASP
met Gly S		40		145	115	
_					r Phe Ala Gli	
Leu Leu A		y Glu Ile	1160	ווו אפ	1165	
	1155			1		د واب
-		r Ser Ala		er Lys se	r Val Gin Ly:	. 417
	1170		1175		1180	
Glu Leu S	Ser Arg Se				r His Leu Al	ยนเก
1185		119			95	
Gly Tyr	Arg Arg G	y Ala Lys	Lys Leu G	lu Ser Se	r Glu Glu As	
1200		1205		1210		1215
Ser Ser	Glu Asp G	lu Glu Lev	Pro Cys P	he Gln Hi	s Leu Leu Ph	e Gly
		220		1225		230
Lvs Val	Asn Asn I	le Pro Ser	Gin Ser 1	ihr Arg Hi	is Ser Thr Va	al Ala
2,5 101	1235		1240		1245	
The Civ		er Lve Acr		Glu Asn Le	eu Leu Ser Le	eu Lys
INT GLU		-, -,5 mai	1255		1260	
	1250			val tie te	eu Ala Lys Al	ls Ser
				44. 11E F.	275	
1265		121			- : -	au Pha
Gln Glu	His His L		u Glu Thr		er Ala Ser Lo	129
1280		1285		1290		
Ser Ser	CID CVC S	er Glu Le	u Glu Asp	Leu Thr A	LB ASN THE A	sn Thr
	GIII Lys 3				4'	
	13	00		1305		310
Gln Asp	13	00				
Gln Asp	13	00		Lys Gln H	et Arg His G 1325	
	13 Pro Phe L 1315	i00 .eu Ile Gl	y Ser Ser 1320	Lys Gin M	et Arg His G	ln Ser

Glu Glu Arg	Gly Thr			u Asn As		Glu Gln Ser
1345		135			1355	
Het Asp Ser	Asn Leu		ALE AL	a Ser Gl	y Cys Glu	Ser Glu Thr
1360		1365			7 0	1375
Ser Val Ser	Glu Asp	Cys Ser	Gly Le	eu Ser Se	r Gln Ser	Asp Ile Leu
	138			1385		1390
Thr Thr Gln	Gln Arg	Asp The	Met G	ln His As	n Leu Ile	Lys Leu Gln
	1395		. 14	400		1405
Gln Glu Met	Ala Giu	Leu Gii	ALB VI	al Leu Gi	u Gin His	Gly Ser Gln
141	0		1415		142	0
Pro Ser Asn	Ser Tyr	Pro Sei	lle I	le Ser As	p Ser Ser	Ala Leu Glu
1425		143	50		1435	
Asp Leu Arg	Asn Pro	Glu Gli	n Ser Ti	hr Ser Gl	u Lys Val	Leu Gln Thr
1440	144	5		1450		1455
Ser Gln Lys	Ser Ser	- Glu Ty	r Pro 1	le Ser Gl	n Asn Pro	Glu Gly Xaa
	1460		14	465		1470
Ser Ala Asp	Lys Phe	e Glu Va	l Ser A	la Asp Se	er Ser Thr	Ser Lys Asn
147	5		1480		148	15
Lys Glu Pro	Gly Va	L Glu Ar	g Ser S	er Pro Se	er Lys Cys	Pro Ser Leu
1490		14	95		1500	
Asp Asp Arg	Trp Ty	r Met Hi	s Ser C	ys Ser G	ly Ser Leu	Gin Asn Arg
1505		1510		1	515	1520
Asn Tyr Pro	Pro Gli	n Glu Gl	u Leu I	le Lys Vi	al Val Asp	Val Glu Glu
	15	25		1530		1535
Gin Gin Leu	Glu Gl	u Ser Gl	y Pro H	is Asp L	eu Thr Glo	u Thr Ser Tyr
	1540		1	545		1550
Leu Pro Arg	Gln As	p Leu Gl	u Gly T	hr Pro T	yr Leu Gla	u Ser Gly Ile
155	i5		1560		15	65
Ser Leu Phe	Ser As	p Asp Pr	o Glu S	er Asp P	ro Ser Gl	u Asp Arg Ala
1570		15	75		1580	
Pro Glu Sei	- Ala Ar	g Val Gl	y Asn I	le Pro S	er Ser Th	r Ser Ala Leu
1585		1590		1	59 5	1600
Lys Val Pro	Gln Le	u Lys Ve	l Ala C	ilu Ser A	la Gln Se	r Pro Ala Ala
	16	05		1610		1615
Ala His Th	r Thr As	p Thr Al	a Gly 1	Tyr Asn A	la Met Gl	u Glu Ser Val
	1620		•	1625		1630
Ser Arg Gl	u Lys Pr	o Glu Le	eu Thr /	Ala Ser T	hr Glu Ar	g Val Asn Lys
. 16			1640			45
Arg Met Se	r Met Va	l Val S	er Gly i	Leu Thr F	ro Glu Gi	u Phe Met Leu
1650			555		1660	
Val Tyr Ly	s Phe Al	a Arg L	s His	His Ile 1	hr Leu Th	r Asn Leu 1le
1665		1670			1675	1680
Thr Glu Gl	u Thr Ti	nr His V	al Val	Met Lys	Thr Asp Al	a Glu Phe Val
		585		1690		1695
Cys Glu Ar	g Thr Lo	eu Lys T	yr Phe	Leu Gly	le Ala Gi	y Gly Lys Trp
	1700	•		1705		1710
Val Val Se	r Tyr Pl	ne Trp V	al Thr	Gin Ser	ile Lys G	lu Arg Lys Met
					17	

Leu	Asn	Glu	His	Asp	Phe	Glu	Val	Arg	Gly	Asp	Val	Val	Asn	Gly	Arg
	1730			1735				1740							
Asn	His	Gln	Gly	Pro	Lys	Arg	Ala	Arg	Glu	Ser	Gln	Asp	Arg	Lys	He
174	5				1750	D				175	5				1760
Phe	Arg	Gly	Leu	Glu	Ile	Cys	Cys	Туг	Gly	Pro	Phe	Thr	Asn	Met	Pro
	_			1765					1770				1775		
Thr	Asp	Gin	Leu	Glu	Trp	Met	Val	Gln	Leu	Cys	Gly	Ala	Ser	Val	Val
		1780										1790			
Lys	Glu	Leu	Ser	Ser	Phe	Thr	Leu	Gly	Thr	Gly	Val	His	Pro	He	Val
•	Arg Glu Lo Val G 1810 Glu Ho	1795			1800			1805							
Val	Val	Gln	Pro	Asp	Ala	Trp	Tht	Glu	Asp	Asn	Gly	Phe	His	Ala	lle
						181					182				
Gly	Gln	Het	Cys	Glu	Ala	Pro	Val	Val	Thr	Arg	Glu	Trp	· Val	Leu	Asp
182	5				183	0				183	5				184
Ser	Val	Ala	Leu	Туг	Gln	Cys	Gln	Glu	Leu	Asp	Thr	Tyr	Leu	ile	Pro
				184	5				185	0				185	i5
Glr	ille	Pro	His	Ser	His	Tyr									
			184												

CLAIMS

What I claim is:

- 1. A method for detecting differential expression of at least one marker gene in pre-invasive cancerous breast tissue, said method comprising the steps of:
- (a) obtaining an abnormal breast tissue sample by a collection step wherein said abnormal breast tissue sample comprises substantially exclusively abnormal breast tissue which exhibits histological or cytological characteristics of pre-invasive breast cancer;
 - (b) isolating mRNA from said abnormal breast tissue sample;
- (c) preparing at least one abnormal breast tissue cDNA library from said mRNA isolated from said abnormal breast tissue sample;
- (d) obtaining a normal breast tissue sample from humans either with or without disease, said normal breast tissue sample comprising substantially exclusively normal breast tissue which does not exhibit histological or cytological characteristics of pre-invasive breast cancer;
- (e) preparing at least one normal breast tissue cDNA library from said normal breast tissue sample; and
- (f) comparing said abnormal breast tissue cDNA library with said normal tissue cDNA library to determine whether the expression of at least one marker gene in said abnormal breast tissue sample is different from the expression of said marker gene in said normal breast tissue sample.
- 2. The method according to Claim 1 wherein said collection step is microscopically-directed.
- 3. The method according to Claim 2 wherein the size of said abnormal tissue sample substantially conforms to an isolatable tissue structure such that only cells exhibiting abnormal cytological or histological characteristics are collected.
- 4. The method according to Claim 3 wherein said isolatable tissue structure comprises ductal epithelial cells in pre-invasive breast cancer tissue.
- 5. The method according to Claim 1 further comprising confirming said differential expression of said marker gene in said normal tissue sample and in said abnormal tissue sample by using a hybridization or PCR technique.

- 6. The method according to Claim 5 wherein said hybridization technique comprises RT-PCR.
- 7. The method according to Claim 5 wherein said hybridization technique comprises nuclease protection assays.
- 8. The method according to Claim 5 wherein said hybridization technique comprises in-situ hybridization of RNA in said abnormal tissue sample and in said normal tissue sample.
- 9. The method according to Claim 1 wherein said abnormal cDNA library and said normal cDNA library are compared by means of differential display.
- 10. The method according to Claim 1 wherein said abnormal cDNA library and said normal cDNA library are compared by means of differential screening.
- 11. The method according to claim 1, wherein said normal tissue comprises normal breast tissue cells.
- 12. The method according to claim 1, wherein said abnormal breast tissue cells are non-comedo ductal carcinoma in situ cells.
- 13. The method according to claim 1, wherein the primer used in the PCR amplification technique is selected from the group consisting of randomly selected primers having the sequences

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5'-CGCGACGGCCGGCGCGTCTGCCAGGG-3', 5'-CTTGCGCGCATACGCACAAC-3',
5'-AACCCTCACCCTAACCCCAA-3', 5'-CGCCCCTGCGTTACCCTCCCGCGG-3',
5'-GGATGGCGTCCTGTAACCCGACGGT-3', 5'-ACTGGGCTGTCCTGCGGTGGCGGGGG-3',
5'-CTGAGAGGTAGCCGCGGGAGGCTG-3', 5'-GCCTGGCCGGACACGGATTACCGC-3',
5'-TTAGCGCATGGTGGACCTGGAGACG-3', 5'-TGTGGTTACGTCAGCGAAGGTAATA-3',
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- 15. A method of diagnosing the presence of pre-invasive breast cancer in human pathologic tissues, said method comprising the steps of:
- (a) obtaining an abnormal breast tissue sample by a collection step wherein said abnormal breast tissue sample comprises substantially exclusively abnormal breast tissue which exhibits histological or cytological characteristics of pre-invasive breast cancer;
 - (b) isolating mRNA from said abnormal breast tissue sample;
- (c) preparing at least one abnormal breast tissue cDNA library from said mRNA isolated from said abnormal breast tissue sample;
- (d) obtaining a normal breast tissue sample from humans either with or without disease, said normal breast tissue sample comprising substantially exclusively normal breast tissue which does not exhibit histological or cytological characteristics of pre-invasive breast cancer;
- (e) preparing at least one normal breast tissue cDNA library from said normal breast tissue sample; and
- (f) comparing said abnormal breast tissue cDNA library with said normal tissue cDNA library to determine whether the expression of at least one marker gene in said abnormal breast tissue sample is different from the expression of said marker gene in said normal breast tissue sample.
- (g) cloning said differentially expressed marker gene using sequence-based amplification to create a cloned marker gene;
 - (h) sequencing said cloned marker gene;
 - (i) producing proteins encoded by said cloned marker gene;

- 18. The method according to claim 15, wherein said medical diagnostic tests comprise blood tests.
- 19. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences designed to clone marker genes encoding secreted proteins.
- 20. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences designed to clone marker genes encoding transcription factors.
- 21. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences designed to clone marker genes encoding RibRed.
- 22. The method according to claim 15, wherein said cloned marker genes encoding secreted proteins are employed in the diagnosis of specific diseases by using a blood test.
- 23. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences adapted to clone marker genes which encode cell surface proteins.
- 24. The method according to claim 15, wherein said proteins encoded by said cloned marker comprise cell surface proteins and wherein the presence of said proteins as a diagnostic indicator is detected by using a diagnostic imaging test.
- 25. A diagnostic method to determine the presence of pre-invasive breast cancer using detection of a differentially expressed marker gene, according to claim 15, wherein said diagnostic method comprises:
- a) obtaining a substantially purified marker gene which is expressed to a greater degree in cells collected by a microscopically-directed cloning method from abnormal tissue than in cells collected from normal tissue;
- b) probing tissues using a hybridization technique to determine whether said substantially purified marker gene is differentially expressed; and,
- c) probing nucleic acids of tissues using a standard hybidization technique to determine the presence of said substantially purified marker gene in a tissue, the

presence of the marker gene indicating the presence of non-comedo DCIS which is preinvasive breast cancer.

26. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:1, which comprises

TTGGGAATTG GGTACGCGGG CCCCCCACTG TGCCGAATTC CTGCATGCGG GGGATCCACT 60
AGTTCAGAGC AGGCCGCCAC CCGTAGGACT CCAGCTTTTG TTCGTTCCCT TTAGTGAGGG 120
TTAATTTTCG AGCTTGGCGT AATCATGGTC ATAGCTGTTT CCTGTGTGAA ATTGTTATCC 180
GCTCACAATT CCACACAACA TACGAGCCGG AAGCATAAAA GTGTAAAGCC TGGGGTGCCT 240
AATGAGTGAG CTAACTCACA TTAA 264

27. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:2, which comprises

TAGCCCGGTT ATCGAAATAG CCACAGCGCC TCTTCACTAT CAGCAGTACG CCGCCCAGTT 60
GTACGGACAC GGA 73

- 28. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:3, which comprises
- TGCCCGATGT GTGTCGTACA ACTGGCGCTG TGGCTGATTT CGATAA 46
- 29. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:4, which comprises

TAGCECATGA GTTCGTGTCC GTACAACTGG GGCGCTGTGG CTGATTTCGA TANNNNNAGC 60
ATCAGCCCGA CG 72

30. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:5, which comprises

TAGCCCGGTT ATCGAAATCA GCCACAGCGC CTAACTTCTG CAGAAGCCTT TGACCATCAC 60
CAGTTGTACG GACACGAACT CATC 84

31. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:6, which comprises

GTGGTTTCCG AAATTCCTGG GAAGGGGGGT GCTGGCGTGT GGAATTGTCG CGGCCCCTGG 60
TCTGCCGCGG CGTTTTTTGT CTACATTCGT CGTAGCTCG 99

32. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:7, which comprises

ATCAGCGCGC GACATICGGG TACCCGCGCC CCCCCCTCCG TCGGAATTCC TCGAGCCGGG 60
ATCCATAGGA TGTGGAGTTA GTTTTGTT 88

- 33. A method for detecting differential expression of at least one marker gene in pre-invasive cancerous breast tissue, said method comprising the steps of:
- (a) obtaining an abnormal tissue sample by a collection step wherein said abnormal tissue sample comprises substantially exclusively abnormal tissue which exhibits histological or cytological characteristics of pre-invasive cancer;
 - (b) isolating mRNA from said abnormal tissue sample;
- (c) preparing at least one abnormal tissue cDNA library from said mRNA isolated from said abnormal tissue sample;
- (d) obtaining a normal tissue sample from humans either with or without disease, said normal tissue sample comprising substantially exclusively normal tissue which does not exhibit histological or cytological characteristics of pre-invasive cancer;
- (e) preparing at least one normal tissue cDNA library from said normal tissue sample; and
- (f) comparing said abnormal tissue cDNA library with said normal tissue cDNA library to determine whether the expression of at least one marker gene in said abnormal tissue sample is different from the expression of said marker gene in said normal tissue sample.
- 34. The method according to Claim 33 wherein said collection step is microscopically-directed.
- a) obtaining a substantially purified marker gene which is expressed to a greater degree in cells collected by a microscopically-directed cloning method from abnormal tissue than in cells collected from normal tissue;
- b) probing tissues using a hybridization technique to determine whether the marker gene is differentially expressed; and,
- c) probing nucleic acids of tissues using a standard PCR technique to determine the presence of the marker gene in a tissue, the presence of the marker gene indicating the presence of pre-invasive cancer.

- 35. Substantially purified DNA having the nucleotide sequences selected from the group of sequences consisting of: SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7.
- 36. An expression vector for the differentially expressed polypeptides encoded by said substantially purified DNA comprising one of the group of DNA sequences of claim 28 operatively linked to at least one control sequence compatible with a suitable bacterial host cell.
- 37. The vector of claim 36 wherein the DNA encoding the differentially expressed polypeptides encoded by said substantially purified DNA comprising one of the group of DNA sequences of claim 28 is linked to at least one sequence from bacteriophage.
- 38. Substantially purified polypeptides encoded by substantially purified DNA comprising one of the group of DNA sequences of claim 35 free of proteins other than proteins encoded by said substantially purified DNA.
- 39. An antibody specifically binding one of the group of polypeptides encoded by one of the nucleotide sequences selected from the group of sequences consisting of: SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID ID NO:7.
- 40. A method of detecting and/or determining said antibody in a test sample, comprising the steps:
 - (a) providing a test sample suspected of containing said marker protein;
- (b) adding a quantity of said marker protein of claim 38 to the antibody of claim 39; and
 - (c) determining a level of said marker protein in said test sample.
- 41. A method of screening compounds for activity in the treatment of breast cancer, comprising the steps of:
 - (a) ligating a DNA sequence that regulates expression of the BRCA1 gene into a vector, the vector having a reporter gene, so that the DNA sequence is located such that the DNA sequence regulates expression of the reporter gene;

- (b) introducing the ligated DNA sequence/reporter gene into a breast cancer cell;
- (c) administering a compound to the breast cancer cell; and
- (d) detecting levels of a protein produced by the reporter cell.
- 42. The method according to claim 41 wherein the DNA sequence is as essentially set forth in SEQ ID NO:48.
- 43. The method according to claim 42 wherein the DNA sequence is selected from among:
 - a. a DNA sequence which hybridizes to SEQ ID NO:48 or fragments thereof; and
 - b. DNA sequences which but for the degeneracy of the genetic code would hybridize to the DNA sequences defined in (a) and (b).
- 44. The method according to claim 41 wherein the ligated DNA sequence/reporter gene is introduced into the breast cancer cell by cloning the ligated DNA sequence/reporter gene into an expression vector and transfecting the breast cancer cells with the expression vector.
- 45. The method according to claim 44 wherein the DNA sequence is essentially set forth in SEQ ID NO:48 or its complementary strands.
 - 46. A method of producing an indicator compound, comprising the steps of:
 - (a) ligating a DNA sequence that regulates expression of the BRCA1 gene into a vector, the vector having a reporter gene, so that the DNA sequence is located such that the DNA sequence regulates expression of the reporter gene;
 - (b) introducing the ligated DNA sequence/reporter gene into a breast cancer cell;
 - (c) administering a biological agent to the breast cancer cell; and
 - (d) producing a protein encoded by the reporter gene; and
 - (e) reacting the protein encoded by the reporter gene with a compound in the reaction media to produce the indicator compound.
- 47. The method according to claim 46 wherein the ligated DNA sequence/reporter gene is introduced into the breast cancer cell by cloning the ligated

DNA sequence/reporter gene into an expression vector and transfecting the breast cancer cells with the expression vector.

- 48. The method according to claim 46 wherein the DNA sequence is as essentially set forth in SEQ ID NO:48 or its complementary strands.
- 49. The method according to claim 46 wherein the DNA sequence is selected from among:
 - a. a DNA sequence which hybridizes to SEQ ID NO:48 or fragments thereof; and
 - b. DNA sequences which but for the degeneracy of the genetic code would hybridize to the DNA sequences defined in (a) and (b).
- 50. A method of treating breast cancer in a patient comprising the steps of ligating a gene that encodes a protein having an amino acid sequence as essentially set forth in SEQ ID NO:49 with a promoter capable of inducing expression of the gene in a breast cancer cell and introducing the ligated gene into a breast cancer cell.
- 51. The method of treating breast cancer described in claim 50 wherein the gene has a DNA sequence selected from among:
 - a. the DNA sequence as essentially set forth in SEQ ID NO:47 or its complementary strands;
 - b. a DNA sequence which hybridizes to SEQ ID NO:47 or fragments thereof; and
 - c. DNA sequences which but for the degeneracy of the genetic code would hybridize to the DNA sequences defined in (a) and (b).
- 52. The method of treating breast cancer described in claim 50 wherein the gene has a DNA sequence having 20-99% homology with SEQ ID NO:47.
- 53. The method according to claim 50 wherein the ligated gene is introduced into the cell in a viral expression vector.
- 54. The method according to claim 50 wherein the breast cancer is genelinked hereditary breast cancer.
- 55. The method described in claim 50 wherein the breast cancer is sporadic breast cancer.

WO 95/19369 PCT/US95/00608

AMENDED CLAIMS

[received by the International Bureau on 14 June 1995 (14.06.95); original claims 13 and 15 amended; new claims 14,16 and 17 added; remaining claims unchanged (8 pages)]

- 6. The method according to Claim 5 wherein said hybridization technique comprises RT-PCR.
- 7. The method according to Claim 5 wherein said hybridization technique comprises nuclease protection assays.
- 8. The method according to Claim 5 wherein said hybridization technique comprises in-situ hybridization of RNA in said abnormal tissue sample and in said normal tissue sample.
- 9. The method according to Claim 1 wherein said abnormal cDNA library and said normal cDNA library are compared by means of differential display.
- 10. The method according to Claim 1 wherein said abnormal cDNA library and said normal cDNA library are compared by means of differential screening.
- 11. The method according to claim 1, wherein said normal tissue comprises normal breast tissue cells.
- 12. The method according to claim 1, wherein said abnormal breast tissue cells are non-comedo ductal carcinoma in situ cells.
- 13. The method according to claim 1, wherein the primer used in the PCR amplification technique is selected from the group consisting of randomly selected primers having the sequences

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20
                5'-CGCGACGGCCGCGCGTCTGCCAGGG-3', 5'-CTTGCGCGCATACGCACAAC-3',
                5'-AACCCTCACCCTAACCCCAA-3', 5'-CGCCCCTGCGTTACCCTCCCCGCCG-3',
                5'-GGATGGCGTCCTGTAACCCGACGCT-3', 5'-ACTGGGCTGTCCTGCGGTGGCGGGG-3',
                5'-CTGAGAGGTAGCCGCGCGGAGGCTG-3', 5'-GCCTGGCCGCGACACGGATTACCGC-3',
                5'-TTAGCGCATGGTGGACCTGGAGACG-3', 5'-TGTGGTTACGTCAGCGAAGGTAATA-3',
                5'-AGTCGCACGCATGTCACGCTCCGCC-3', 5'-TATCCAAGCGGCAGGCTACGAGGCC-3',
25
                5'-GGCGCGCCCGACGGTCTGGTATCTA-3', 5'-CTCCCTCCCCGGACTCGGGGTTAGT-3',
                5'-ATGCGGGCGGCTCGGGCCTGGTCGC-3', 5'-CGTGAAGCCTATGCCCTCCCTCAAC-3',
                5'-GTGCCGTCGTAGCCCTTCAGCGATC-3', 5'-GCGACACTAGGCTCCCGGAGGAGGG-3',
                5'-TGGGCCAGGCCTCCGGGCCCGGTAT-3', 5'-CCGGAACTGCGATAGCGTCCGTCCC-3',
                5'-AGCGGACACCTGTTTCCCGAGAGCC-3', 5'-AACGGGTGGACATCCGCCTGCCGCC-3',
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                5'-TGAACCACGATGTCAATCGTCCCGA-3', 5'-TCATCCCCGCCGAAAGACGCTCGCC-3',
                5'-ATAGGCTGCGGCACGCGCTGGGACT-3', 5'-GACCAGGTGCGCACGAGCATGTACA-3',
                5'-AGCGTAGTCATCGGCCTTCGCGCCC-3', 5'-GGCCCCTAGCCCAGGGTGAAGCCCA-3',
                5'-CCCAGTGCTACGGGCCGCCCCAAGC-3', 5'-CCTTCCTGGGTTACCTGCCCTCGGG-3',
                5'-TCCGGACAGCAGCCACGCCAAGGGC-3', 5'-ACGCGCTGGTCCACCGAGGCCTGAT-3',
 35
                5'-CGATGCAAGGCCAGCAGCACTCGAC-3', 5'-CCCCCGGAGCGGACCACCGGACGTG-3',
                5'-AGCGGGGAGGGATCGGGGGCCAAGC-3', 5'-GCCTGGTGTAGGCAGGCAGCTCTTA-3',
                5'-CCACCCCTGTAGTGCGGGGTGCGAG-3', 5'-GGAACCCGACGCCCGTCCAGGGTTC-3',
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5'-TCGGGCAGCAAGGCCGGGACGCTCC-3', 5'-GACGGGGGACGGGCTAGGTGGCTTA-3', and 5'-CTTGTTGCCGGCGAGAGGGCTGCC-3'.

- 14. The method according to claim 2, wherein said abnormal tissue sample is approximately 2 mm in diameter.
- 15. A method of diagnosing the presence of pre-invasive breast cancer in human pathologic tissues, said method comprising the steps of:
- (a) obtaining an abnormal breast tissue sample by a collection step wherein said abnormal breast tissue sample comprises substantially exclusively abnormal breast tissue which exhibits histological or cytological characteristics of pre-invasive breast cancer;
 - (b) isolating mRNA from said abnormal breast tissue sample;
- (c) preparing at least one abnormal breast tissue cDNA library from said mRNA isolated from said abnormal breast tissue sample;
- (d) obtaining a normal breast tissue sample from humans either with or without disease, said normal breast tissue sample comprising substantially exclusively normal breast tissue which does not exhibit histological or cytological characteristics of pre-invasive breast cancer;
- (e) preparing at least one normal breast tissue cDNA library from said normal breast tissue sample; and
- (f) comparing said abnormal breast tissue cDNA library with said normal tissue cDNA library to determine whether the expression of at least one marker gene in said abnormal breast tissue sample is different from the expression of said marker gene in said normal breast tissue sample.
- (g) cloning said differentially expressed marker gene using sequence-based amplification to create a cloned marker gene;
 - (h) sequencing said cloned marker gene;
 - (i) producing proteins encoded by said cloned marker gene;
- (j) generating antibodies which will recognize said proteins encoded by said cloned marker gene by antigen recognition; and
 - (k) detecting said recognized antigen by means of medical diagnostic tests.
- 16. The method according to claim 15, wherein said medical diagnostic tests comprise diagnostic tissue tests.
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- 17. The method according to claim 15, wherein said medical diagnostic tests comprise X-ray tests.
- 18. The method according to claim 15, wherein said medical diagnostic tests comprise blood tests.
- 19. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences designed to clone marker genes encoding secreted proteins.
- 20. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences designed to clone marker genes encoding transcription factors.
- 21. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences designed to clone marker genes encoding RibRed.
- 22. The method according to claim 15, wherein said cloned marker genes encoding secreted proteins are employed in the diagnosis of specific diseases by using a blood test.
- 23. The method according to claim 15, wherein said sequence-based amplification employs DNA sequences adapted to clone marker genes which encode cell surface proteins.
- 24. The method according to claim 15, wherein said proteins encoded by said cloned marker comprise cell surface proteins and wherein the presence of said proteins as a diagnostic indicator is detected by using a diagnostic imaging test.
- 25. A diagnostic method to determine the presence of pre-invasive breast cancer using detection of a differentially expressed marker gene, according to claim 15, wherein said diagnostic method comprises:
- a) obtaining a substantially purified marker gene which is expressed to a greater degree in cells collected by a microscopically-directed cloning method from abnormal tissue than in cells collected from normal tissue;
- b) probing tissues using a hybridization technique to determine whether said substantially purified marker gene is differentially expressed; and,
 - c) probing nucleic acids of tissues using a standard hybidization technique

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to determine the presence of said substantially purified marker gene in a tissue, the presence of the marker gene indicating the presence of non-comedo DCIS which is pre-invasive breast cancer.

26. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:1, which comprises

TIGGGAATIG GGTACGCGGG CCCCCCACTG TGCCGAATIC CTGCATGCGG GGGATCCACT 60
AGTTCAGAGC AGGCCGCCAC CCGTAGGACT CCAGCTTTIG TTCGTTCCCT TTAGTGAGGG 120
TTAATTTTCG AGCTTGGCGT AATCATGGTC ATAGCTGTTT CCTGTGTGAA ATTGTTATCC 180
GCTCACAATI CCACACAACA TACGAGCCGG AAGCATAAAA GTGTAAAGCC TGGGGTGCCT 240
AATGAGTGAG CTAACTCACA TTAA 264

- 27. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:2, which comprises TAGCCCGGTT ATCGAAATAG CCACAGCGCC TCTTCACTAT CAGCAGTACG CCGCCCAGTT 60 GTACGGACAC GGA
- 28. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:3, which comprises

TGCCCGATGT GTGTCGTACA ACTGGCGCTG TGGCTGATTT CGATAA 46

29. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:4, which comprises

TAGCCCATGA GTTCGTGTCC GTACAACTGG GGCGCTGTGG CTGATTTCGA TANNNHNAGC 60
ATCAGCCCGA CG 72

30. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:5, which comprises

TAGCCCGGTT ATCGAAATCA GCCACAGCGC CTAACTTCTG CAGAAGCCTT TGACCATCAC 60
CAGTTGTACG GACACGAACT CATC 84

31. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:6, which comprises

GTGGTTTCCG AAATTCCTGG GAAGGGGGGT GCTGGCGTGT GGAATTGTCG CGGCCCCTGG 60
TCTGCCGCGG CGTTTTTTGT CTACATTCGT CGTAGCTCG 99

32. The method according to claim 25, wherein said substantially purified marker gene has the sequence listed according to SEQ ID NO:7, which comprises

ATCAGCGCGC GACATTCGGG TACCCGCGCC CCCCCCTCCG TCGGAATTCC TCGAGCCGGG 60
ATCCATAGGA TGTGGAGTTA GTTTTGTT 88

33. A method for detecting differential expression of at least one marker gene in pre-invasive cancerous breast tissue, said method comprising the steps of:

- (a) obtaining an abnormal tissue sample by a collection step wherein said abnormal tissue sample comprises substantially exclusively abnormal tissue which exhibits histological or cytological characteristics of pre-invasive cancer;
 - (b) isolating mRNA from said abnormal tissue sample;
- (c) preparing at least one abnormal tissue cDNA library from said mRNA isolated from said abnormal tissue sample;
- (d) obtaining a normal tissue sample from humans either with or without disease, said normal tissue sample comprising substantially exclusively normal tissue which does not exhibit histological or cytological characteristics of pre-invasive cancer;
- (e) preparing at least one normal tissue cDNA library from said normal tissue sample; and
- (f) comparing said abnormal tissue cDNA library with said normal tissue cDNA library to determine whether the expression of at least one marker gene in said abnormal tissue sample is different from the expression of said marker gene in said normal tissue sample.
- 34. The method according to Claim 33 wherein said collection step is microscopically-directed.
- a) obtaining a substantially purified marker gene which is expressed to a greater degree in cells collected by a microscopically-directed cloning method from abnormal tissue than in cells collected from normal tissue;
- b) probing tissues using a hybridization technique to determine whether the marker gene is differentially expressed; and,
- c) probing nucleic acids of tissues using a standard PCR technique to determine the presence of the marker gene in a tissue, the presence of the marker gene indicating the presence of pre-invasive cancer.

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- 35. Substantially purified DNA having the nucleotide sequences selected from the group of sequences consisting of: SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEO ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID NO:7.
- 36. An expression vector for the differentially expressed polypeptides encoded by said substantially purified DNA comprising one of the group of DNA sequences of claim 28 operatively linked to at least one control sequence compatible with a suitable bacterial host cell.
- 37. The vector of claim 36 wherein the DNA encoding the differentially expressed polypeptides encoded by said substantially purified DNA comprising one of the group of DNA sequences of claim 28 is linked to at least one sequence from bacteriophage.
- 38. Substantially purified polypeptides encoded by substantially purified DNA comprising one of the group of DNA sequences of claim 35 free of proteins other than proteins encoded by said substantially purified DNA.
- 39. An antibody specifically binding one of the group of polypeptides encoded by one of the nucleotide sequences selected from the group of sequences consisting of: SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, and SEQ ID ID NO:7.
- 40. A method of detecting and/or determining said antibody in a test sample, comprising the steps:
 - (a) providing a test sample suspected of containing said marker protein;
- (b) adding a quantity of said marker protein of claim 38 to the antibody of claim 39; and
 - (c) determining a level of said marker protein in said test sample.
- 41. A method of screening compounds for activity in the treatment of breast cancer, comprising the steps of:
 - (a) ligating a DNA sequence that regulates expression of the BRCA1 gene into a vector, the vector having a reporter gene, so that the DNA sequence is located such that the DNA sequence regulates expression of the reporter gene;
 - (b) introducing the ligated DNA sequence/reporter gene into a breast cancer

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cell;

- (c) administering a compound to the breast cancer cell; and
- (d) detecting levels of a protein produced by the reporter cell.
- 42. The method according to claim 41 wherein the DNA sequence is as essentially set forth in SEQ ID NO:48.
- 43. The method according to claim 42 wherein the DNA sequence is selected from among:
 - a. a DNA sequence which hybridizes to SEQ ID NO:48 or fragments thereof; and
 - b. DNA sequences which but for the degeneracy of the genetic code would hybridize to the DNA sequences defined in (a) and (b).
- 44. The method according to claim 41 wherein the ligated DNA sequence/reporter gene is introduced into the breast cancer cell by cloning the ligated DNA sequence/reporter gene into an expression vector and transfecting the breast cancer cells with the expression vector.
- 45. The method according to claim 44 wherein the DNA sequence is essentially set forth in SEQ ID NO:48 or its complementary strands.
 - 46. A method of producing an indicator compound, comprising the steps of:
 - (a) ligating a DNA sequence that regulates expression of the BRCA1 gene into a vector, the vector having a reporter gene, so that the DNA sequence is located such that the DNA sequence regulates expression of the reporter gene;
 - (b) introducing the ligated DNA sequence/reporter gene into a breast cancer cell:
 - (c) administering a biological agent to the breast cancer cell; and
 - (d) producing a protein encoded by the reporter gene; and
 - (e) reacting the protein encoded by the reporter gene with a compound in the reaction media to produce the indicator compound.
- 47. The method according to claim 46 wherein the ligated DNA sequence/reporter gene is introduced into the breast cancer cell by cloning the ligated DNA sequence/reporter gene into an expression vector and transfecting the breast

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cancer cells with the expression vector.

- 48. The method according to claim 46 wherein the DNA sequence is as essentially set forth in SEQ ID NO:48 or its complementary strands.
- 49. The method according to claim 46 wherein the DNA sequence is selected from among:
 - a. a DNA sequence which hybridizes to SEQ ID NO:48 or fragments thereof; and
 - b. DNA sequences which but for the degeneracy of the genetic code would hybridize to the DNA sequences defined in (a) and (b).
- 50. A method of treating breast cancer in a patient comprising the steps of ligating a gene that encodes a protein having an amino acid sequence as essentially set forth in SEQ ID NO:49 with a promoter capable of inducing expression of the gene in a breast cancer cell and introducing the ligated gene into a breast cancer cell.
- 51. The method of treating breast cancer described in claim 50 wherein the gene has a DNA sequence selected from among:
 - a. the DNA sequence as essentially set forth in SEQ ID NO:47 or its complementary strands;
 - b. a DNA sequence which hybridizes to SEQ ID NO:47 or fragments thereof; and
 - c. DNA sequences which but for the degeneracy of the genetic code would hybridize to the DNA sequences defined in (a) and (b).
- 52. The method of treating breast cancer described in claim 50 wherein the gene has a DNA sequence having 20-99% homology with SEQ ID NO:47.
- 53. The method according to claim 50 wherein the ligated gene is introduced into the cell in a viral expression vector.
- 54. The method according to claim 50 wherein the breast cancer is genelinked hereditary breast cancer.
- 55. The method described in claim 50 wherein the breast cancer is sporadic breast cancer.

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STATEMENT UNDER ARTICLE 19

Pursuant to Article 19 of the Patent Cooperation Treaty and Rule 46, Applicant respectfully submits the attached sheets of amended claims. The sheets are replacement sheets for pages 98-105 of the above referenced International application. These sheets contain Claims 6-55 of the above referenced international application. New Claims 14, 16 and 17 have been added to replacement pages 99-100. Additional primers have been listed in Claim 13 on replacement pages 98-99. These primers are described in the Sequence Listing. Claim 15 has been amended to include steps (j) and (k) on replacement page 99. The new claims and the amended claims do not go beyond the scope of the application as filed. The remaining replacement sheets include no amendments, but are filed to maintain the correct numbering of the claim pages.

Figure 1:

Anatomic Lesion Types in the Human Breast with Pre-malignant Implication TABLE I:

(Dupont, et al, 1985 and 1993.) (Page, et al, 1991.) Reference < .00001 > < .00001 P value Relative Risk* 9-10 fold 4-5 fold Determinant Lesions with Regional Risk Indicators of generalized increased risk Atypical ductal hyperplasia Pre-malignant Lesions Lobular CIS

* represents the 95% confidence interval for relative risk.

(Page, et al, 1982.)

< .00005

10-11 fold

Non-comedo DCIS

Figure 1: Table I describes anatomic lesion types in the human breast with pre-malignant implication.

Fig. 2

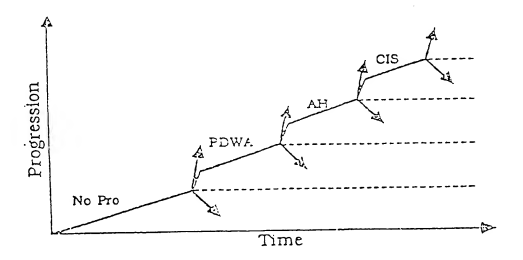


Fig. 3

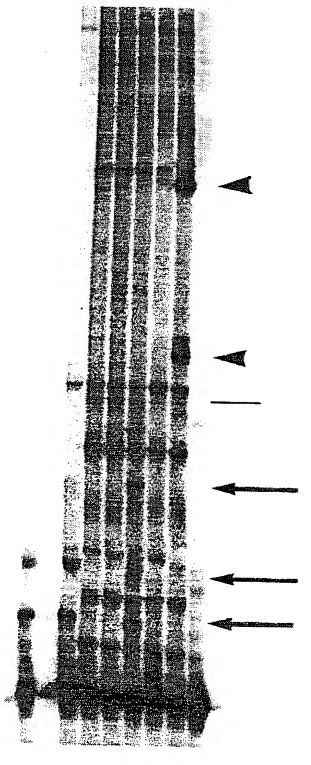


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Fig. 3



Fig. 5



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Figure 6: Comparison of the sequence between DCIS-1 and the human and hamster genes.

Fig. 7

Con NL1 NL2 NL3 #12 #6 #4 #6 #10 #100





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Fig. 8 - Table of the Genetic Code

Amino Acids			Codons					
Alanine	Ala	A	GCA	GCC	GCG	ecn		
Cysteine	Cys	c	ugc	UGU				•
Aspartic acid	Asp	D	GAC	GAU				
Glutamic acid	Glu	E	AAD	GAG				
Phenylalanine	Phe	F	UUC	UUU				
Glycine	Gly	G	GGA	GGC	ccc	GGU		
Histidine	His	н	CAC	CAU				
Isoleucine	Ile	1	AUA	SUA	AUU			
Lysine	Lys	ĸ	AAA	AAG				
Leucine	Leu	L	AUU	UUG	CUA	cuc	cuc	CUU
Methionine	Met	M	AUG					
Asparagine	Asn	И	AAC	UAA				
Proline	Prc	P	433	ccc	ccc	ccu		
Glutamine	Gln	0	AAD	CAG				
Arginine	Arg	R	AGA	AGG	CGA	CGC	CGG	CGU
Serine	Ser	\$	ASC	AGU	UCA	UCC	ucc	UCU
Threonine	Thr	ĭ	AC#	ACC	ACG	ACU		
Valine	Val	v	4U2	GUC	GUG	GUU		
Tryptophan	1rp	¥	იმი					
Tyrosine	lyr	Y	DAU	UAU				

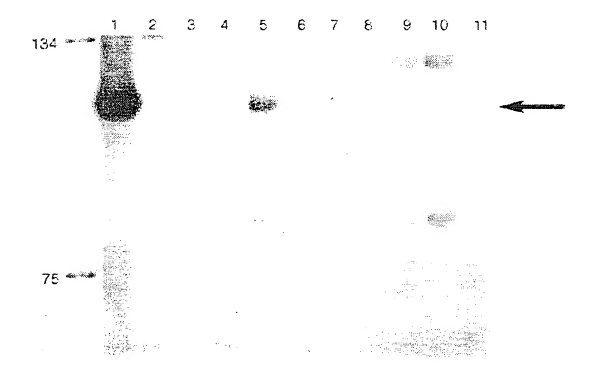
Figure 8: Table of the Genetic Code.

SEQ ID NO): 1: (DCIS-1)				
TIGGGAATIG GGT	ACGCGGG CCCCCCACTG	TGCCGAATTC	CTGCATGCGG	GGGATCCACT	60
AGTTCAGAGC CCG	TAGGACT CCAGCTTTTG	TICGIICCCT	TTAGTGAGGG	TTAATTITCG	120
AGCTTGGCGT AAT	CATGGTC ATCCTGTGTG	AAATTGTTAT	CCGCTCACAA	TTCCACACAA	180
CATACGAGEE GGA	AGCATAA AAGTGTAAGC	AATGAGTGAG	CTAACTCACA	AATT	234
SEO ID NO: 2:	(DC15-2)				
TAGCCEGGTT ATC	GAAATAG CCACAGEGEE	TCTTCACTAT	CAGCAGTACG	CCGCCCAGTT	60
GTA CGG ACA CG	iG				72
SEQ 10 HO: 3:					
TGCCCGATGA GTT	IGTGTCGT ACAACTGGCG	CTGTGGCTGA	TTTCGATAA		45
SEQ ID NO: 4:					
TAGCCCATGA GTT	ICGIGICC GTACAACIGG	GGCGCTGTGG	CTGATTTCGA	TAHNNNHAGC	60
ATCAGCCCGA CG					72
SEO 10 NO: 5:					
	CGAAATCA GCCACAGCGC	CTAACTICTG	CAGAAGCCTT	TGACCATCAC	60
CAGTIGIACE GAA	AACGAACT CATC				84
SEO 10 NO: 6:					
	ATTCCTG GGAAGGGGGG			GEGGCCCTG	60
GTCTGCCGCG GC	GITITIT GICTACATIC	GTCGTAGCTC	С		10
SEG 10 NO: 7:					
	CATTEGGG TACCEGEGE		TEGGAATTE	TEGAGEEGG	60 88
ATTATAGGA TG	TGGAGTTA GTTTTGTT				08

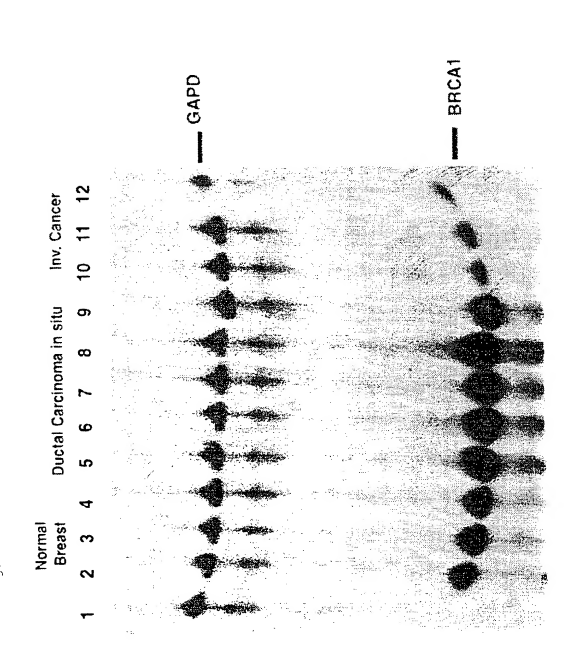
Figure 9: Table of Differentially Expressed Marker Genes From Pre-Invasive Human Breast Tissue

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Fig. 10A



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Fig. 11A

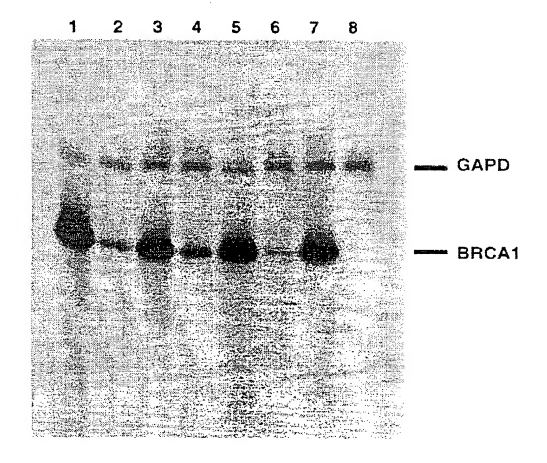
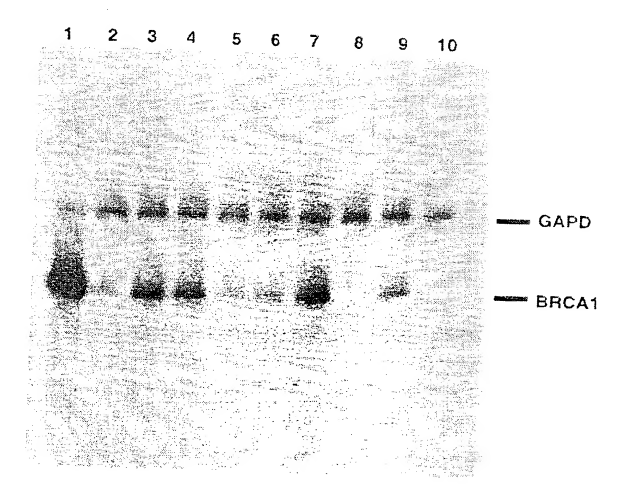


Fig. 11B



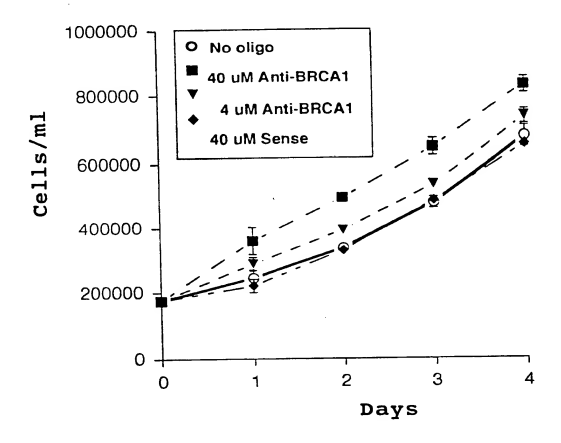


Fig. 12A

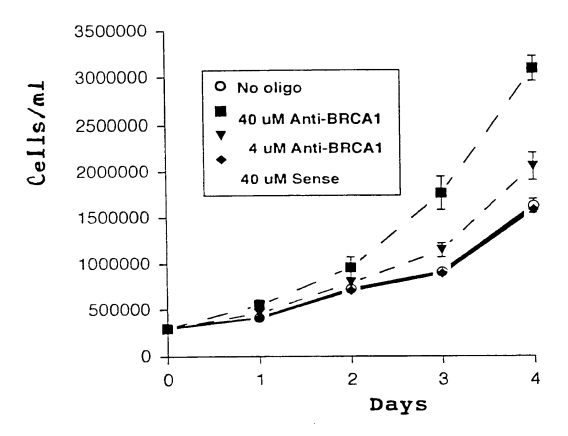


Fig. 12B

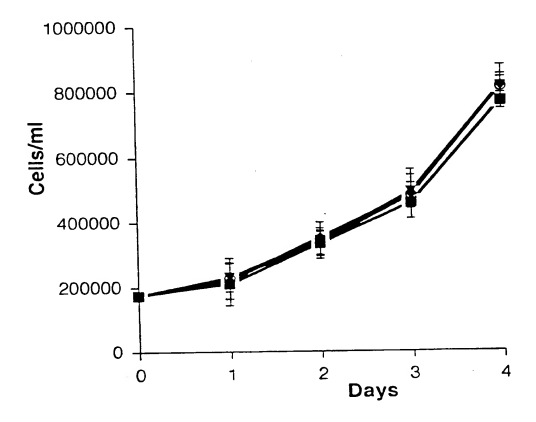


Fig. 12C

Fig. 13A

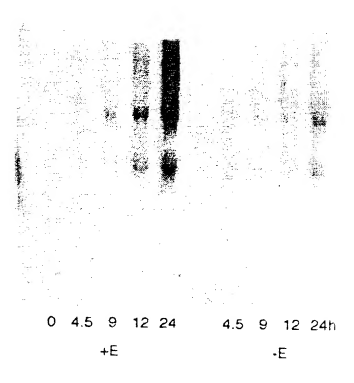
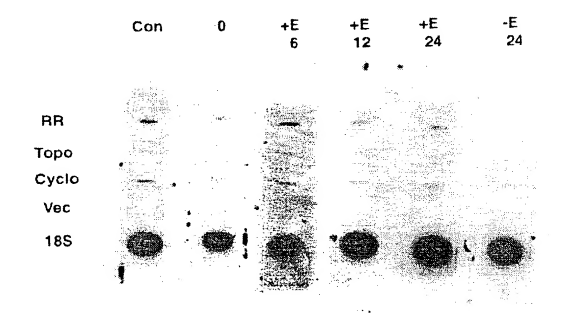
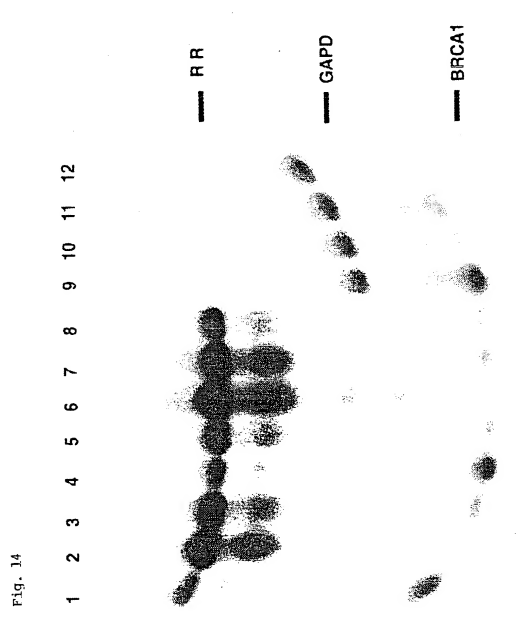


Fig. 13B





SUBSTITUTE SHEET (RULE 26)

Inc. al application No. PCT/US95/00608

IPC(6) :Please See Extra Sheet.								
According to	US CL: Please See Extra Sheet. According to International Patent Classification (IPC) or to both national classification and IPC							
B. FIEL	DS SEARCHED							
	cumentation searched (classification system followed b							
	35/6, 7.1, 69.2, 172.3, 320.1; 514/44; 536/23.2, 23	<u> </u>						
Documentati	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched							
Electronic da	ata base consulted during the international search (nam	e of data base and, where practicable,	search terms used)					
	ee Extra Sheet.							
C. DOC	UMENTS CONSIDERED TO BE RELEVANT							
Category*	Citation of document, with indication, where appr	ropriate, of the relevant passages	Relevant to claim No.					
P, Y	Science, Vol. 266, issued 07 Octobra and Strong Candidate for the Bread Susceptibility Gene BRCA1", padocument.	1-13, 15, 18- 37, 40						
Y	Cancer Research, Vol. 52, issued Liang et al., "Differential Display an RNAs from Breast Cancer <i>versus</i> M pages 6966-6968, see entire docu	1-13, 15, 18-37						
Y	Cancer Surveys, Vol. 18, issued "Histopathology: Old Principles ar 1-16, Tables 1 and 2 and Figures 1	1-13						
	(D-6	See patent family annex.						
	her documents are listed in the continuation of Box C.	*T hater document published after the in	ternational filing date or priority					
1	pecial categories of cited documents: ocument defining the general state of the art which is not considered	date and not in conflict with the appli principle or theory underlying the in	cation but cited to understand the					
to to	be of particular relevance	*Y* document of particular relevance;	he claimed invention cannot be					
1	*E* cartier document published on or after the international filing date considered novel or cannot be considered to involve an inventive step when the document is taken alone.							
cited to establish the publication date of another citation or other special reason (as specified) considered to involve an inventive step when the document								
n	ocument referring to an oral disclosure, use, exhibition or other ocass ocument published prior to the international filing date but later than	being obvious to a person skilled in the art document member of the same patent family						
the priority date claimed								
Date of the	e actual completion of the international search	04 MAY 1995						
Commiss Box PCT	ioner of Patents and Trademarks	Stephanie W. Zitomer, Ph.D.						
Washingt	on, D.C. 20231 No. (703) 305-3230	Telephone No. (703) 308-0196						

Form PCT/ISA/210 (second sheet)(July 1992)*

nal application No.
PCT/US95/00608

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Y	DNA (N.Y.), Vol. 5, No. 5, issued 1986, Neuhold et al., "Dioxin-Inducible Enhancer Region Upstream from the Mouse P-1450 Gene and Interaction with a Heterologous SV-40 Promoter", abstract, see entire document.	41-49
E, Y	US, A, 5,399,346 (ANDERSON ET AL.) 21 March 1995.	50-55

Form PCT/ISA/210 (continuation of second sheet)(July 1992)*

in . . . nal application No. PCT/US95/00608

Box 1 Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 1-13, 15, 18-37, 40-55
No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.

Form PCT/ISA/210 (continuation of first sheet(1))(July 1992)+

International application No. PCT/US95/00608

A. CLASSIFICATION OF SUBJECT MATTER:

IPC (6):

C07H 21/02, 21/04; C12Q 1/68; G01N 33/53; C12P 21/00; C12N 15/63, 15/85; A61K 48/00

A. CLASSIFICATION OF SUBJECT MATTER:

US CL :

435/6, 7.1, 69.2, 172.3, 320.1; 514/44; 536/23.2, 23.5; 935/3, 6, 9, 11, 14, 23, 77, 78

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, DIALOG: Biosis, Derwent Biotech. Abstracts, WPI, Chem. Abstr., Diss. Abstr., Embase, Medline, Current Biotech. Abstr. (Royal Soc.); search strat: (cancer or carcinoma)(p)(breast or ovar?)(p)(gene or nucleic)(p)gene(..) sequence, nucleic(..) sequence?

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING

This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group 1, claims 1-13 and 25-37, drawn to a first process of using and a first product used: a diagnostic process using the product nucreic acids.

Group II, claims 15, 18-24 and 40, drawn to a second process of using, an immunoassay.

Group III, claims 38 and 39, drawn to a second product used, proteins including polypeptides and antibodies.

Group IV, claims 41-45, drawn to a third process of using, a process of screening compounds for activity in breast cancer treatment.

Group V, claims 46-49, drawn to a fourth process of using, a process for producing an indicator compound.

Group VI, claims 50-55, drawn to a fifth process of using, a process for treating breast cancer.

The inventions listed as Groups I-VI do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

The inventions of Groups III-VI lack the diagnostic technical feature and the inventions of Groups II-VI lack the nucleic acid special feature of Group I while the inventions of Groups II and IV-VI each have a different result such that they individually lack the special features of the others that are responsible for that result: The Group II process has a diagnostic result; the Group IV invention identifies a compound that affects the expression of the BRCA1 gene; the Group V process produces an indicator compound; the invention of Group VI treats cancer.

Form PCT/ISA/210 (extra sheet)(July 1992)+

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